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medicament in a mammalian host wherein the gene constru	e conju	geting moiety and a heterologous prodrug activating enzyme for use as a upable of expressing the cell targeting moiety and enzyme as a conjugate gate is directed to leave the cell thereafter for selective localisation at a
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#### CHEMICAL COMPOUNDS

This invention relates particularly to gene directed enzyme prodrug therapy (GDEPT) using *in situ* antibody generation to provide enhanced selectivity, particularly for use in cancer 5 therapy.

Known gene therapy based prodrug therapeutic approaches include virus-directed enzyme prodrug therapy (VDEPT) and gene-directed enzyme prodrug therapy (GDEPT), the latter term encompassing both VDEPT and non-viral delivery systems. VDEPT involves targeting tumour cells with a viral vector carrying a gene which codes for an enzyme capable of activating a prodrug. The viral vector enters the tumour cell and enzyme is expressed from the enzyme gene inside the cell. In GDEPT, alternative approaches such as microinjection, liposomal delivery and receptor mediated DNA uptake as well as viruses may be used to deliver the gene encoding the enzyme.

In both VDEPT and GDEPT the enzyme gene can be transcriptionally regulated by

15 DNA sequences capable of being selectively activated in mammalian cells e.g. tumour cells

(EP 415 731 (Wellcome); Huber et al, Proc. Natl. Acad. Sci. USA, 88, 8039-8043,1991).

While giving some degree of selectivity, gene expression may also occur in non-target cells

and this is clearly undesirable when the approach is being used to activate prodrugs into

potent cytotoxic agents. In addition these regulatory sequences will generally lead to reduced

20 expression of the enzyme compared with using viral promoters and this will lead to a reduced

ability to convert prodrug in the target tissue.

Expression and localisation of the prodrug activating enzyme inside the cell has disadvantages. Prodrug design is severely limited by the fact that the prodrug has to be able to cross the cell membrane and enter the cell but not be toxic until it is converted to the drug 25 inside the cell by the activating enzyme. Most prodrugs utilise hydrophilic groups to prevent cell entry and thus reduce cytotoxicity. Prodrug turnover by activating enzyme produces a less hydrophilic drug which can enter cells to produce anti-cancer effects. This approach can not be used when the activating enzyme is expressed inside the cell. Another disadvantage is that target cells which lack intracellular activating enzyme will be difficult to attack because 30 they are unable to generate active drug. To achieve this desirable "bystander activity" (or "neighbouring cell kill"), the active drug will have to be capable of diffusion out of the cell containing activating enzyme to reach target cells which lack enzyme expression. Many

active drugs when produced inside a cell will be unable to escape from the cell to achieve this bystander effect.

Modifications of GDEPT have been put forward to overcome some of the problems described above. Firstly vectors have been described which are said to express the activating 5 enzyme on the surface of the target cell (WO 96/03515) by attaching a signal peptide and transmembrane domain to the activating enzyme. The approach, if viable, would overcome the problems of having the activating enzyme located inside the cell but would still have to rely on transcriptionally regulated sequences capable of being selectively expressed in target cells to restrict cell expression. As described above there are disadvantages of using such 10 sequences. Secondly vectors have been described which result in secretion of the enzyme from the target cell (WO 96/16179). In this approach the enzyme would be able to diffuse away from its site of generation since it is extracellular and not attached to the cell surface. Enzyme which has diffused away from the target site would be capable of activating prodrug at non-target sites leading to unwanted toxicity. To achieve some selectivity it is suggested 15 that enzyme precursors could be used which are cleaved by pathology associated proteases to form active enzyme. Some selectivity is likely to be achieved by this approach but its unlikely that activation will only occur at target sites. In addition, once activated, the enzyme will still be free to diffuse away from the target site and thus suffer from the same drawback described above.

For GDEPT approaches, three levels of selectivity can be observed. Firstly, there is selectivity at the cell infection stage such that only specific cell types are targeted. For example cell selectivity can be provided by the gene delivery system per se. An example of this type of selectivity is set out in International Patent Application WO 95/26412 (UAB Research Foundation) which describes the use of modified adenovirus fiber proteins incorporating cell specific ligands. Other examples of cell specific targeting include ex vivo gene transfer to specific cell populations such as lymphocytes and direct injection of DNA into muscle tissue.

The second level of selectivity is control of gene expression after cell infection such as for example by the use of cell or tissue specific promoters. If the gene has been delivered to a 30 cell type in a selective manner then it is important that a promoter is chosen that is compatible with activity in the cell type.

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The third level of selectivity can be considered as the selectivity of the expressed gene construct. Selectivity at this level has received scant attention to date. In International patent application WO 96/16179 (Wellcome Foundation) it is suggested that enzyme precursors could be used which are cleaved by pathology associated proteases to form active enzyme.

5 Some selectivity is likely to be achieved by this approach but it is unlikely that activation will only occur at target sites. In addition, once activated, the enzyme will still be free to diffuse away from the target site and thus suffer from the same drawback of activating prodrug at non-target sites leading to unwanted toxicity.

There exists a need for more selective GDEPT systems to reduce undesirable effects in 10 normal tissues arising from erroneous prodrug activation.

The present invention is based on the discovery that antibody-heterologous enzyme gene constructs can be expressed intracellularly and used in GDEPT systems (or other systems such as AMIRACS - see below) for cell targeting arising from antibody specificity to deliver cell surface available enzyme in a selective manner. This approach may be used optionally in combination with any other suitable specificity enhancing technique(s) such as targeted cell infection and/or tissue specific expression.

According to one aspect of the present invention there is provided a gene construct encoding a cell targeting antibody and a heterologous enzyme for use as a medicament in a mammalian host wherein the gene construct is capable of expressing the antibody and enzyme 20 as a conjugate within a target cell in the mammalian host and wherein the conjugate can leave the cell thereafter for selective localisation at a cell surface antigen recognised by the antibody.

According to another aspect of the present invention there is provided a gene construct encoding a cell targeting moiety and a heterologous prodrug activating enzyme for use as a medicament in a mammalian host wherein the gene construct is capable of expressing the cell targeting moiety and heterologous prodrug activating enzyme as a conjugate within a cell in the mammalian host and wherein the conjugate is directed to leave the cell thereafter for selective localisation at a cell surface antigen recognised by the cell targeting moiety.

The "cell targeting moiety" is defined as any polypeptide or fragment thereof which selectively binds to a particular cell type in a host through recognition of a cell surface antigen. Preferably the cell targeting moiety is an antibody. Cell targeting moieties other than antibodies include ligands as described for use in Ligand Directed Enzyme Prodrug

Therapy as described in International patent application WO 97/26918, Cancer Research Campaign Technology Limited, such as for example epidermal growth factor, heregulin, cerbB2 and vascular endothelial growth factor with the latter being preferred.

A "cell targeting antibody" is defined as an antibody or fragment thereof which

5 selectively binds to a particular cell type in a host through recognition of a cell surface antigen. Preferred cell targeting antibodies are specific for solid tumours, more preferably colorectal tumours, more preferably an anti-CEA antibody, more preferably antibody A5B7 or 806.077 antibody with 806.077 antibody being especially preferred. Hybridoma 806.077 antibody was deposited at the European Collection of Animal Cell Cultures (ECACC), PHLS

10 Centre for Applied Microbiology & Research, Porton Down, Salisbury, Wiltshire SP4 0JG, United Kingdom on 29th February 1996 under accession no. 96022936 in accordance with the Budapest Treaty.

Antibody A5B7 binds to human carcinoembryonic antigen (CEA) and is particularly suitable for targeting colorectal carcinoma. A5B7 is available from DAKO Ltd., 16 Manor

15 Courtyard, Hughenden Avenue, High Wycombe, Bucks HP13 5RE, England, United Kingdom. In general the antibody (or antibody fragment) - enzyme conjugate should be at least divalent, that is to say capable of binding to at least 2 tumour associated antigens (which may be the same or different). Antibody molecules may be humanised by known methods such as for example by "CDR grafting" as disclosed in EP239400 or by grafting complete variable regions from for example a murine antibody onto human constant regions ("chimaeric antibodies") as disclosed in US 4816567. Humanised antibodies may be useful for reducing immunogenicity of an antibody (or antibody fragment). A humanised version of antibody A5B7 has been disclosed in International Patent Application WO 92/01059 (Celltech).

The hybridoma which produces monoclonal antibody A5B7 was deposited with the European Collection of Animal Cell Cultures, Division of Biologics, PHLS Centre for Applied Microbiology and Research, Porton Down, Salisbury, Wiltshire SP4 OJG, United Kingdom. The date of deposit was 14th July 1993 and the accession number is No. 93071411. Antibody A5B7 may be obtained from the deposited hybridoma using standard techniques known in the art such as documented in Fenge C, Fraune E & Schuegerl K in "Production of Biologicals from Animal Cells in Culture" (Spier RE, Griffiths JR & Meignier B, eds) Butterworth-Heinemann, 1991, 262-265 and Anderson BL & Gruenberg ML

in "Commercial Production of Monoclonal Antibodies" (Seaver S, ed), Marcel Dekker, 1987, 175-195. The cells may require re-cloning from time to time by limiting dilution in order to maintain good levels of antibody production.

A "heterologous enzyme" is defined as an enzyme for turning over a substrate that has

been administered to the host and the enzyme is not naturally present in the relevant
compartment of the host. The enzyme may be foreign to the mammalian host (e.g. a bacterial
enzyme like CPG2) or it may not naturally occur within the relevant host compartment (e.g.
the use of lysozyme as an ADEPT enzyme (for an explanation of ADEPT see below) is
possible because lysozyme does not occur naturally in the circulation, see US 5433955, Akzo

NV). The relevant host compartment is that part of the mammalian host in which the
substrate is distributed. Preferred enzymes are enzymes suitable for ADEPT or AMIRACS
(Antimetabolite with Inactivation of Rescue Agents at Cancer Sites; see Bagshawe (1994) in
Cell Biophysics 24/25, 83-91) but ADEPT enzymes are preferred. Antibody directed enzyme

15 selective antibody conjugated to an enzyme. The conjugate is administered to the patient (usually intravenously), allowed to localise at the tumour site(s) and clear from the blood and other normal tissues. A prodrug is then administered to the patient which is converted by the enzyme (localised at the tumour site) into a cytotoxic drug which kills the tumour cells.

prodrug therapy (ADEPT) is a known cancer therapeutic approach. ADEPT uses a tumour

In International Patent Application WO 96/20011, published 4-Jul-96, we proposed a 20 "reversed polarity" ADEPT system based on mutant human enzymes having the advantage of low immunogenicity compared with for example bacterial enzymes. A particular host enzyme was human pancreatic CPB (see for example, Example 15 [D253K]human CPB & 16 [D253R]human CPB therein) and prodrugs therefor (see Examples 18 & 19 therein). The host enzyme is mutated to give a change in mode of interaction between enzyme and prodrug in

- 25 terms of recognition of substrate compared with the native host enzyme. In our subsequent International Patent Application No PCT/GB96/01975 (published 6-Mar-97 as WO 97/07796) further work on mutant CPB enzyme/ prodrug combinations for ADEPT are described. Preferred enzymes suitable for ADEPT are any one of CPG2 or a reversed polarity CPB enzyme, for example any one of [D253K]HCPB, [G251T,D253K]HCPB or
- 30 [A248S,G251T,D253K]HCPB. A preferred form of CPG2 is one in which the polypeptide glycosylation sites have been mutated so as to prevent or reduce glycosylation on expression in mammalian cells (see WO 96/03515, Cancer Research Campaign Technology); this gives

Guthrie and G R Fink.

improved enzyme activity. Further considerations arise for enzymes such as CPB which require a pro domain to facilitate correct folding; here the pro domain can either be expressed as a separately (in trans) or expressed as part of the fusion protein and subsequently removed.

Large scale purification of CPG2 from *Pseudomonas* RS-16 was described in

5 Sherwood *et al* (1985), Eur, J. Biochem., <u>148</u>, 447 - 453. CPG2 may be obtained from Centre for Applied Microbiology and Research, Porton Down, Salisbury, Wiltshire SP4 0JG, United Kingdom. CPG2 may also be obtained by recombinant techniques. The nucleotide coding sequence for CPG2 has been published by Minton, N.P. *et al.*, Gene, <u>31</u> (1984), 31-38. Expression of the coding sequence has been reported in *E.coli* (Chambers, S.P. <u>et al.</u>, Appl.

- Microbiol, Biotechnol. (1988), 29, 572-578) and in Saccharomyces cerevisiae (Clarke, L. E. et al., J. Gen Microbiol, (1985) 131, 897-904). Total gene synthesis has been described by M. Edwards in Am. Biotech. Lab (1987), 5, 38-44. Expression of heterologous proteins in E.coli has been reviewed by F.A.O. Marston in DNA Cloning Vol. III, Practical Approach Series, IRL Press (Editor D M Glover), 1987, 59-88. Expression of proteins in yeast has
  been reviewed in Methods in Enzymology Volume 194, Academic Press 1991, Edited by C.
- Whilst cancer therapeutic approaches are preferred the invention may also be applied to other therapeutic areas as long as a target antigen can be selected and a suitable enzyme/ prodrug combination prepared. For example, inflammatory diseases such as rheumatiod arthritis may be treated by for example using an antibody selective for synovial cells fused to an enzyme capable of converting an anti-inflammatory drug in the form of a prodrug into an anti-inflammatory drug. Use of antibodies to target rheumatoid arthritis disease has been described in Blakey et al, 1988, Scand. J. Rheumatology, Suppl. 76, 279-287.

A "conjugate" between antibody and enzyme can be a fusion protein (covalent 25 linkage) or the conjugate can be formed by non-covalent binding between antibody and enzyme formed in situ. Preferably the conjugate is in the form of a fusion protein, more preferably the antibody component of the fusion is at least divalent (for improved binding avidity compared with monovalent antibody). Antibody constructs lacking an Fc portion are preferred, especially Fab or F(ab')<sub>2</sub> fragments. For CPG2 fusions (or fusions with any non-monomeric enzyme) special considerations apply because CPG2 is a dimeric enzyme and the antibody is preferably divalent thus there exists the potential for undesirable competing dimerisation between two molecular species. Therefore a preferred CPG2 fusion is one in

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which the fusion protein is formed through linking a C-terminus of an antibody Fab heavy chain (ie lacking a hinge region) to an N-terminus of a CPG2 molecule; two of these Fab-CPG2 molecules then dimerise through the CPG2 dimerisation domain to form a (Fab-CPG2)2 conjugate. For antibody constructs with monomeric enzymes, F(ab')2 fragments are 5 preferred, especially F(ab')<sub>2</sub> fragments having a human IgG3 hinge region. Fusions between antibody and enzyme may optionally be effected through a short peptide linker such as for example (G<sub>4</sub>S)<sub>3</sub>. Preferred fusion constructs are those in which the enzyme is fused to the C terminus of the antibody, through the heavy or light chain thereof with fusion through the antibody heavy chain being preferred. Accordingly a preferred gene construct is a gene 10 construct for use as a medicament as described herein in which the antibody-enzyme CPG2 conjugate is a fusion protein in which the enzyme is fused to the C terminus of the antibody through the heavy or light chain thereof whereby dimerisation of the encoded conjugate when expressed can take place through a dimerisation domain on CPG2. A more preferred gene construct is a gene construct for use as a medicament wherein the fusion protein is formed 15 through linking a C-terminus of an antibody Fab heavy chain to an N-terminus of a CPG2 molecule to form a Fab-CPG2 whereby two Fab-CPG2 molecules when expressed dimerise through CPG2 to form a (Fab-CPG2)2 conjugate. In another embodiment of the invention a preferred gene construct for use as a medicament is one wherein the carboxypeptidase is selected from [D253K]HCPB, [G251T,D253K]HCPB or [A248S,G251T,D253K]HCPB.

It is contemplated that should it be possible to obtain a natural multimeric enzyme in monomeric form whilst substantially retaining enzymic activity then the monomeric form of the enzyme could be used to form a conjugate of the invention. Similarly, it is contemplated that should it be possible to obtain a natural monomeric enzyme in multimeric form whilst substantially retaining enzymic activity then the multimeric form of the enzyme could be used to form a conjugate of the invention.

The conjugate is directed to leave the cell after expression therein through use of a secretory leader sequence which is cleaved as the conjugate passes through the cell membrane. Preferably the secretory leader is the secretory leader that occurs naturally with the antibody.

According to another aspect of the present invention there is provided use of a gene construct encoding a cell targeting antibody and a heterologous enzyme for use for manufacture of a medicament for cancer therapy in a mammalian host wherein the gene

construct is capable of expressing the antibody and enzyme as a conjugate within a target cell in the mammalian host and wherein the conjugate can leave the cell thereafter for selective localisation at a cell surface antigen recognised by the antibody.

Any suitable delivery system may be applied to deliver the gene construct of the present invention including viral and non-viral systems. Viral systems include retroviral vectors, adenoviral vectors, adeno-associated virus, vaccinia, herpes simplex virus, HIV, the minute virus of mice, hepatitis B virus and influenza virus. Non-viral systems include uncomplexed DNA, DNA-liposome complexes, DNA-protein complexes and DNA-coated gold particles.

Retroviral vectors lack immunogenic proteins and there is no preexisting host immunity but are limited to infecting dividing cells. Retroviruses have been used in clinical trials (Rosenberg et al., N. Engl. J. Med., 1990, 323: 570-578). Retroviruses are composed of an RNA genome that is packaged in an envelope derived from host cell membrane and viral proteins. For gene expression, it must first reverse transcribe its positive-strand RNA genome into double-stranded DNA, which is then integrated into the host cell DNA using reverse transcriptase and integrase protein contained in the retrovirus particle. The integrated provirus is able to use host cell machinery for gene expression.

Murine leukemia virus is widely used (Miller et al., Methods Enzymol., 1993, 217: 581-599). Retroviral vectors are constructed by removal of the gag, pol and env genes to 20 make room for the relevant payload and to eliminate the replicative functions of the virus. Virally encoded mRNAs are eliminated and this removes any potential immune response to the transduced cells. Genes encoding antibiotic resistance often are included as a means of selection. Promoter and enhancer functions also may be included for example to provide for tissue-specific expression after administration in vivo. Promoter and enhancer functions 25 contained in the long terminal repeat may also be used.

These viruses can be produced only in viral packaging cell lines. The packaging cell line may be constructed by stably inserting the deleted viral genes (gag, pol. and env) into the cell such that they reside on different chromosomes to prevent recombination. The packaging cell line is used to construct a producer cell line that will generate replication-defective retrovirus containing the relevant payload gene by inserting the recombinant proviral DNA. Plasmid DNA containing the long terminal repeat sequences flanking a small portion of the gag gene that contains the encapsidation sequence and the genes of interest is transfected into

the packaging cell line using standard techniques for DNA transfer and uptake (electroporation, calcium precipitation, etc.). Variants of this approach have been employed to decrease the likelihood of production of replication-competent virus (Jolly, D., Cancer Gene Therapy, 1994, 1, 51-64). The host cell range of the virus is determined by the envelope gene (env) and substitution of env genes with different cell specificities can be employed. Incorporation of appropriate ligands into the envelope protein may also be used for targeting.

Administration may be achieved by any suitable technique e.g. ex vivo transduction of patients' cells, by the direct injection of virus into tissue, and by the administration of the 10 retroviral producer cells.

The ex vivo approach has a disadvantage in that it requires the isolation and maintenance in tissue culture of the patient's cells, but it has the advantage that the extent of gene transfer can be quantified readily and a specific population of cells can be targeted. In addition, a high ratio of viral particles to target cells can be achieved and thus improve the transduction efficiency (Anderson et al., Hum. Gene Ther., 1990, 1: 331-341; Rosenberg et al., N. Engl. J. Med., 1990, 323: 570-578; Culver et al., Hum. Gene Ther., 1991, 2: 107-109Nienhuis et al., Cancer, 1991, 67: 2700-2704, Anderson et al., Hum. Gene Ther., 1990, 1: 331-341, Grossman et al., Nat. Genet., 1994, 6: 335-341, Lotze et al., Hum. Gene Ther., 1992, 3: 167-177; Lotze, M.T., Cell Transplant., 1993, 2: 33-47; Lotze et al., Hum. Gene Ther., 1994, 5: 41-55 and US patent 5399346 (Anderson). In some cases direct introduction of virus in vivo is necessary. Retroviruses have been used to treat brain turnours wherein the ability of a retrovirus to infect only dividing cells (tumour cells) may be particularly advantageous.

To increase efficiency Oldfield et al., in Hum. Gene Ther., 1993, 4: 39-69 proposed

25 the administration of a retrovirus producer cell line directly into patients' brain tumours. The
murine producer cell would survive within the brain tumour for a period of days, and would
secrete retrovirus capable of transducing the surrounding brain tumour. Virus carrying the
herpes virus thymidine kinase gene renders cells susceptible to killing by ganciclovir, which is
metabolized to a cytotoxic compound by thymidine kinase. Patent references on retroviruses

30 are: EP 334301, WO 91/02805 & WO 92/05266 (Viagene) and; US 4650764 (University of
Wisconsin).

Human adenoviral infections have been described (see Horwitz, M.S., In Virology, 2<sup>nd</sup> ed. Raven Press, New York, 1990, pp. 1723-1740). Most adults have prior exposure to adenovirus and have antiadenovirus antibodies. These viruses possess a double-stranded DNA genome, and replicate independent of host cell division.

Adenoviral vectors possess advantageous properties. They are capable of transducing a broad spectrum of human tissues and high levels of gene expression can be obtained in dividing and nondividing cells. Several routes of administration can be used including intravenous, intrabiliary, intraperitoneal, intravesicular, intracranial and intrathecal injection, and direct injection of the target organ. Thus targeting based on anatomical boundaries is 10 feasible.

The adenoviral genome encodes about 15 proteins and infection involves a fiber protein to bind a cell surface receptor. The penton base of the capsid engages integrin receptor domains (α<sub>3</sub>β<sub>3</sub>, or α<sub>3</sub>β<sub>5</sub>) on the cell surface resulting in internalization of the virus. Viral DNA enters the nucleus and begins transcription without cell division. Expression and replication is under control by the E1A and E1B genes (see Horwitz, M.S., In Virology, 2<sup>nd</sup> ed., 1990, pp. 1723-1740). Removal of E1 genes renders the virus replication-incompetent. Expression of adenoviral proteins leads to both an immune response which may limit effectiveness particularly on repeat administration. However, recent approaches in which other adenoviral genes such as the E2a gene (which controls expression of the fibre knob and a number of other viral proteins) are also removed from the viral genome may abolish or greatly reduce the expression of many of these viral proteins in target cells.

Adenoviral serotypes 2 and 5 have been extensively used for vector construction. Bett et al., Proc. Nat. Acad. Sci. U.S.A., 1994, 91: 8802-8806 have used an adenoviral type 5 vector system with deletions of the E1 and E3 adenoviral genes. The 293 human embryonic 25 kidney cell line has been engineered to express E1 proteins and can thus transcomplement the E1-deficient viral genome. The virus can be isolated from 293 cell media and purified by limited dilution plaque assays (Graham, F.L. and Prevek, L. In Methods in Molecular Biology: Gene Transfer and Expression Protocols, Humana Press 1991, pp. 109-128). Recombinant virus can be grown in 293 cell line cultures and isolated by lysing infected cells and purification by caesium chloride density centrifugation. One problem of the 293 cells for manufacture of recombinant adenovirus is that due to additional flanking regions of the E1 genes is that they may give rise to replication competent adenovirus (RCA) during the viral

particle production. Although this material is only wild type adenovirus and not replication competent recombinant virus it can have significant effects on the eventual yield of the desired adenoviral material and lead to increased manufacturing costs, quality control issues for the production runs and acceptance of batches for clinical use. Alternative cell lines such as the PER.C6 which have more defined E1 gene integration than 293 cells (i.e. contain not flanking viral sequence) have been developed which do not allow the recombination events which produce RCA and thus have the potential to overcome above viral production issues.

Adenoviral vectors have the disadvantage of relatively short duration of transgene expression due to immune system clearance and dilutional loss during target cell division but improvements in vector design are anticipated. Patent references on adenoviruses are: WO 96/03517 (Boehringer); WO 96/13596 (Rhone Poulenc Rorer); WO 95/29993 (University of Michigan) and; WO 96/34969 (Canji). Recent advances in adenoviral vectors for cancer gene therapy including the development of strategies to reduce immunogenicity, chimeric adenoviral/retroviral vectors and conditional (or restricted) replicatiative recombinant adenoviral systems are reviewed in Bilbao et al., Exp. Opin. Ther. Patents, 1997, 7 (12):1427-1446.

Adeno-associated virus (AAV) (Kotin, R.M., Hum. Gene Ther., 1994, <u>5</u>: 793-801) are single-stranded DNA, nonautonomous parvoviruses able to integrate into the genome of nondividing cells of a very broad host range. AAV has not been shown to be associated with 20 human disease and does not elicit an immune response.

AAV has two distinct life cycle phases. Wild-type virus will infect a host cell, integrate and remain latent. In the presence of adenovirus, the lytic phase of the virus is induced, which is dependent on the expression of early adenoviral genes, and leads to active virus replication. The AAV genome is composed of two open reading frames (called rep and cap) flanked by inverted terminal repeat (ITR) sequences. The rep region encodes four proteins which mediate AAV replication, viral DNA transcription, and endonuclease functions used in host genome integration. The rep genes are the only AAV sequences required for viral replication. The cap sequence encodes structural proteins that form the viral capsid. The ITRs contain the viral origins of replication, provide encapsidation signals, and participate in viral DNA integration. Recombinant, replication-defective viruses that have been developed for gene therapy lack rep and cap sequences. Replication-defective AAV can be produced by cotransfecting the separated elements necessary for AAV replication into a

permissive 293 cell line. Patent references on AAV include: WO 94/13788 (University of Pittsburgh) and US 4797368 (US Department of Health).

Gene therapy vectors from pox viruses have been described (Moss, B. and Flexner, C., Annu. Rev. Immunol., 1987, 5: 305-324; Moss, B., In Virology, 1990, pp. 2079-2111).

- 5 Vaccinia are large, enveloped DNA viruses that replicate in the cytoplasm of infected cells.

  Nondividing and dividing cells from many different tissues are infected, and gene expression from a nonintegrated genome is observed. Recombinant virus can be produced by inserting the transgene into a vaccinia-derived plasmid and transfecting this DNA into vaccinia-infected cells where homologous recombination leads to the virus production. A significant
- 10 disadvantage is that it elicits a host immune response to the 150 to 200 virally encoded proteins making repeated administration problematic.

The herpes simplex virus is a large, double-stranded DNA virus that replicates in the nucleus of infected cells suitable for gene delivery (see Kennedy, P.G.E. and Steiner, I., Q.J. Med., 1993, 86: 697-702). Advantages include a broad host cell range, infection of dividing and nondividing cells, and large sequences of foreign DNA can be inserted into the viral genome by homologous recombination. Disadvantages are the difficulty in rendering viral preparations free of replication-competent virus and a potent immune response. Deletion of the viral thymidine kinase gene renders the virus replication-defective in cells with low levels of thymidine kinase. Cells undergoing active cell division (e.g., tumour cells) possess sufficient thymidine kinase activity to allow replication. Cantab Pharmaceuticals have a published patent application on herpes viruses (WO 92/05263).

A variety of other viruses, including HIV, the minute virus of mice, hepatitis B virus, and influenza virus, have been considered as possible vectors for gene transfer (see Jolly, D., Cancer Gene Therapy, 1994, 1: 51-64).

The use of attenuated Salmonella Typhimurium bacteria which specifically target and replicate in hypoxic environments (such as are found in the necrotic centres of tumours) as gene delivery vehicles for prodrug enzyme based therapy (Tumour Amplified Prodrug Enzyme Therapy known as TAPET™) has also been proposed and is under development by Vion Pharmaceuticals. This system offers a further gene delivery alternative to the viral and non-viral delivery approaches discussed below.

Nonviral DNA delivery strategies are also applicable. These DNA delivery systems include uncomplexed plasmid DNA, DNA-liposome complexes, DNA-protein complexes,

and DNA-coated gold particles.

Purified nucleic acid can be injected directly into tissues and results in transient gene expression for example in muscle tissue, particularly effective in regenerating muscle (Wolff et al., Science, 1990, 247: 1465-1468). Davis et al., in Hum. Gene Ther., 1993, 4: 733-740 has published on direct injection of DNA into mature muscle. Skeletal and cardiac muscle is generally preferred. Patent references are: WO 90/11092, US 5589466 (Vical) and WO 97/05185 (biodegradable DNA impregnated hydrogels for injection, Focal).

Plasmid DNA on gold particles can be "fired" into cells (e.g. epidermis or melanoma) using a gene-gun. DNA is coprecipitated onto the gold particle and then fired using an electric spark or pressurized gas as propellant (Fynan *et al.*, Proc. Natl. Acad. Sci. U.S.A., 1993, 90: 11478-11482). Electroporation has also been used to enable transfer of DNA into solid tumours using electroporation probes employing multi-needle arrays and pulsed, rotating electric fields (Nishi *et al.*, in Cancer Res., 1996, 56:1050-1055). High efficiency gene transfer to subcutaneous tumours has been claimed with significant cell transfection enhancement and better distribution characteristics over intra-tumoural injection procedures.

Liposomes work by surrounding hydrophilic molecules with hydrophobic molecules to facilitate cell entry. Liposomes are unilamellar or multilamellar spheres made from lipids. Lipid composition and manufacturing processes affect liposome structure. Other molecules can be incorporated into the lipid membranes. Liposomes can be anionic or cationic.

- 20 Nicolau et al., Proc. Natl. Acad. Sci. U.S.A., 1983, 80: 1068-1072 has published on insulin expression from anionic liposomes injected into rats. Anionic liposomes mainly target the reticuloendothelial cells of the liver, unless otherwise targeted. Molecules can be incorporated into the surface of liposomes to alter their behavior, for example cell-selective delivery (Wu, G.Y. and Wu, C.H., J. Biol. Chem., 1987, 262: 4429-4432).
- Felgner et al., Proc. Nat. Acad. Sci. U.S.A., 1987, <u>84</u>: 7413-7417 has published on cationic liposomes, demonstrated their binding of nucleic acids by electrostatic interactions and shown cell entry. Intravenous injection of cationic liposomes leads to transgene expression in most organs on injection into the afferent blood supply to the organ. Cationic liposomes can be administered by aerosol to target lung epithelium (Brigham et al., Am. J.
- 30 Med. Sci., 1989, <u>298</u>: 278-281). Patent references on liposomes are: WO 90/11092, WO 91/17424, WO 91/16024, WO 93/ 14788 (Vical) and; WO 90/01543 (Intracel).

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In-Vivo studies with cationic liposome transgene delivery have been published by: Nabel et al., Rev. Hum. Gene Ther., 1994, 5: 79-92; Hyde et al., Nature, 1993, 362: 250-255 and; Conary et al., J. Clin. Invest., 1994, 93: 1834-1840).

Microparticles are being studied as systems for delivery of DNA to phagocytic cells

5 such approaches have been pursued by Pangaea Pharmaceuticals in their ENDOSHERE<sup>TM</sup>

DNA microencapsulation delivery system which has been used to effect more efficient transduction of phagocytic cells such as macrophages which ingest the microspheres. The microspheres encapsulate plasmid DNA encoding potentially immunogenic peptides which when expressed lead to peptide display via MHC molecules on the cell surface which can stimulate immune response against such peptides and protein sequences which contain the same epitopes. This approach is presently aimed towards a potential role in anti-tumour and pathogen vaccine development but may have other possible gene therapy applications.

In the same way as synthetic polymers have been used to package DNA natural viral coat proteins which are capable of homogeneous self-assembly into Virus-like particles

15 (VLPs) have been used to package DNA. The major structural coat protein VP1 of human polyoma virus can be expressed as a recombinant protein and is able to package plasmid DNA during self-assembly into a VLP. The resulting particles can be subsequently used to transduce various cell lines, while preliminary studies show little immunogenic response to such VP1 based VLPs. Such systems may offer an attractive intermediate between synthetic polymer non-viral vectors and the alternative viral delivery systems since they may offer combined advantages e.g. simplicity of production and high level transduction efficiency.

To improve the specificity of gene delivery and expression the therapeutic gene the inclusion of targeting elements into the delivery vehicles and the use of regulatory expression elements have been investigated both singlulary and in combination in many of the previously described delivery systems.

Improvements in DNA vectors have also been made and are likely applicable to all of the non-viral delivery systems. These include the use of supercoiled minicircles reported by RPR Gencell (which do not have bacterial origins of replication nor antibiotic resistance genes and thus are potentially safer as they exhibit a high level of biological containment), episomal expression vectors as developed by Copernicus Gene Systems Inc (replicating episomal expression systems where the plasmid amplifies within the nucleus but outside the chromosome and thus avoids genome integration events) and T7 systems as developed by

Progenitor (a strictly a cytoplasmic expression vector in which the vector itself expresses phage T7 RNA polymerase and the therapeutic gene is driven from a second T7 promoter, using the polymerase generated by the first promoter). Other, more general improvements to DNA vector technology include use of cis-acting elements to effect high levels of expression (Vical), sequences derived from alphoid repeat DNA to supply once-per-cell-cycle replication and nuclear targeting sequences (from EBNA-1 gene (Calos at Stanford, with Megabios); SV40 early promoter/enhancer or peptide sequences attached to the DNA).

Targeting systems based on cell receptor recognition by ligand linked to DNA have been described by Michael, S.I. and Curiel, D.T., Gene Therapy, 1994, 1: 223-232. Using the ligand recognized by such a receptor the DNA becomes selectively bound and internalized into the target cell (Wu, G.Y. and Wu, C.H., J. Biol. Chem., 1987, 262: 4429-4432). Poly-Llysine (PLL), a polycation, has been used to couple a variety of protein ligands to DNA by chemical cross-linking methods. DNA is electrostatically bound to PLL-ligand molecules. Targetting systems have been published by Zenke et al., Proc. Nat. Acad. Sci. U.S.A., 1990, 87: 3655-3659 using transferrin receptor; Wu, G.Y. and Wu, C.H., J. Biol. Chem., 1987, 262: 4429-4432 using the asialoorosomucoid receptor, and Batra et al., Gene Therapy, 1994, 1: 255-260, using cell surface carbohydrates. Agents such as chloroquine or co-localised adenovirus can be used to reduce DNA degradation in the lysosomes (see Fisher, K.J. and Wilson, J.M., Biochem. J., 1994, 299, 49-58). Cristiano et al., Proc. Natl. Acad. Sci. U.S.A., 1993, 90: 11548-11552 has constructed adenovirus-DNA-ligand complexes. Patent references on receptor mediated endocytosis are: WO 92/05250 (asialoglycoproteins, University of Connecticut) and US 5354844 (transferrin receptor, Boehringer).

DNA and ligand can be coated over the surface of the adenovirus to create a coated adenovirus (Fisher, K.J. and Wilson, J.M., Biochem. J., 1994, 299, 49-58). However the presence of two receptor pathways for DNA entry (ligand receptor and adenovirus receptor) reduces the specificity of this delivery system but the adenovirus receptor pathway can be eliminated by using an antibody against adenovirus fiber protein as the means for linkage to DNA (Michael, S.I. and Curiel, D.T., Gene Therapy, 1994, 1: 223-232). Use of purified endosomalytic proteins rather than intact adenovirus particles is another option (Seth, P., J. 30 Virol., 1994, 68: 1204-1206).

The expression of a gene construct of the invention at its target site is preferably under the control of a transcriptional regulatory sequence (TRS). A TRS is a promoter optionally

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combined with an enhancer and/or an control element such as a genetic switch described below.

One example of a TRS is a "genetic switch" that may be employed to control expression of a gene construct of the invention once it has been delivered to a target cell. 5 Control of gene expression in higher eucaryotic cells by procaryotic regulatory elements (which are preferred for the present invention) has been reviewed by Gossen et al in TIBS, 18th December 1993, 471-475. Suitable systems include the E.coli lac operon and the especially preferred E.coli tetracycline resistance operon. References on the tetracycline system include Gossen et al (1995) Science 268, 1766; Damke et al (1995) Methods in 10 Enzymology 257, Academic Press; Yin et al (1996) Anal. Biochem. 235, 195 and; patents US 5464758, US 5589362, WO 96/01313 and WO 94/29442 (Bujard). An ecdysone based switch (International Patent Appln No.PCT/GB96/01195, Publication No. WO 96/37609, Zeneca) is another option. Other options are listed below. Connaught Laboratories (WO-93/20218) describe a synthetic inducible eukaryotic promoter comprising at least two different classes of 15 inducible elements. Rhone-Poulenc Rorer (WO 96/30512) describe a tetracycline-related application for a conditional gene expression system. Ariad (WO 94/18317) describes a protein dimerisation based system for which in vivo activity has been shown. Bert O'Malley of the Baylor College of Medicine (WO 93/23431, US 5364791, WO 97/10337) describes a molecular switch based on the use of a modified steroid receptor. The Whitehead Institute 20 have an NF-KB inducible gene expression system (WO 88/05083). Batelle Memorial have described a stress inducible promoter (European patent EP 263908).

Examples of TRSs which are independent of cell type include the following: cytomegalovirus promoter/ enhancer, SV40 promoter/ enhancer and retroviral long terminal repeat promoter/ enhancer. Examples of TRSs which are dependent on cell type (to give an 25 additional degree of targeting) include the following promoters: carcinoembryonic antigen (CEA) for targeting colorectal, lung and breast; alpha-foetoprotein (AFP) for targeting transformed hepatocytes; tyrosine hydroxylase, choline acetyl transferase or neurone specific enolase for targeting neuroblastomas; insulin for targeting pancreas and; glial fibro acidic protein for targeting glioblastomas. Some oncogenes may also be used which are selectively 30 expressed in some tumours e.g. HER-2/neu or c-erbB2 in breast and N-myc in neuroblastoma.

Accordingly, a preferred gene construct for use as a medicament is a construct comprising a transcriptional regulatory sequence which comprises a promoter and a control

element which is a genetic switch to control expression of the gene construct. A preferred genetic switch control element is regulated by presence of tetracycline or ecdysone. A preferred promoter is dependent on cell type and is selected from the following promoters: carcinoembryonic antigen (CEA); alpha-foetoprotein (AFP); tyrosine hydroxylase; choline acetyl transferase; neurone specific enolase; insulin; glial fibro acidic protein; HER-2/neu; cerbB2; and N-myc. Preferably the gene construct for use as a medicament described herein is packaged within an adenovirus for delivery to the mammalian host. A general review of targeted gene therapy is given in Douglas et al., Tumor Targeting, 1995, 1: 67-84.

The antibody encoded by the gene construct of the invention may be any form of
antibody construct such as for example F(ab')<sub>2</sub>; F(ab'), Fab, Fv, single chain Fv & V-min. Any
suitable antibody construct is contemplated, for example a recently described antibody
fragment is "L-F(ab)<sub>2</sub>" as described by Zapata (1995) in Protein Engineering, 8, 1057-1062.

Disulphide bonded Fvs are also contemplated. For constructs based on CPG2 enzyme, Fab
fragment constructs dimerised through enzyme dimerisation are preferred. Non-human
antibodies may be humanised for use in humans to reduce host immune responses. A
humanized antibody, related fragment or antibody binding structure is a polypeptide
composed largely of a structural framework of human derived immunoglobulin sequences
supporting non human derived amino acid sequences in and around the antigen binding site
(complementarity determining regions or CDRs). Appropriate methodology has been
described for example in detail in WO 91/09967, EP 0328404 and Queen et al. Proc Natl
Acad Sci 86,10029, Mountain and Adair (1989) Biotechnology and Genetic Engineering
Reviews 10, 1 (1992) although alternative methods of humanisation are also contemplated
such as antibody veneering of surface residues (EP 519596, Merck/NIH, Padlan et al).

According to another aspect of the present invention there is provided a matched two

25 component system designed for use in a mammalian host in which the components comprise:

(i) a first component that comprises a gene construct encoding a cell targeting antibody and a
heterologous prodrug activating enzyme wherein the gene construct is capable of expressing
the antibody and enzyme as a conjugate within a target cell in the mammalian host and
wherein the conjugate can leave the cell thereafter for selective localisation at a cell surface

30 antigen recognised by the antibody and;

(ii) a second component that comprises a prodrug which can be converted into an active drug by the enzyme.

Antibody directed enzyme prodrug therapy (ADEPT) is a known cancer therapeutic approach. ADEPT uses a tumour selective antibody conjugated to an enzyme. The conjugate is administered to the patient (usually intravenously), allowed to localise at the tumour site(s) and clear from the blood and other normal tissues. A prodrug is then administered to the patient which is converted by the enzyme (localised at the tumour site) into a cytotoxic drug which kills the tumour cells.

The present invention can be applied to any ADEPT system. Suitable examples of ADEPT systems include those based on any of the following enzymes: carboxypeptidase G2; carboxypeptidase A; aminopeptidase; alkaline phosphatase; glycosidases; β-glucuronidase;

- penicillin amidase; β-lactamase; cytosine deaminase; nitroreductase; or mutant host enzymes including carboxypeptidase A, carboxypeptidase B, and ribonuclease. Suitable references on ADEPT systems include Melton RG (1996) in J. National Cancer Institute 88, 1; Niculescu-Duvaz I (1995) in Current Medicinal Chemistry 2, 687; Knox RJ (1995) in Clin. Immunother. 3, 136; WO 88/07378 (CRCT); Blakey et al, Cancer Res. 56, 3287-92, 1996; US 5587161
- 15 (CRCT and Zeneca); WO 97/07769 (Zeneca); and WO 95/13095 (Wellcome). The heterologous enzyme may be in the form of a catalytic antibody; see for example EP 745673 (Zeneca). A review articles on ADEPT systems include Hay & Denny (1996), Drugs of the Future, 21(9), 917-931 and Blakey (1997). Exp. Opin. Ther. Patents, 7(9), 965-977.

A preferred matched two component system is one in which:

- the first component comprises a gene encoding the heterologous enzyme CPG2; and the second component prodrug is selected from N-(4-[N,N-bis(2-iodoethyl)amino]-phenoxycarbonyl)-L-glutamic acid, N-(4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl)-L-glutamic-gamma-(3,5-dicarboxy)anilide or N-(4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl)-L-glutamic acid or a pharmaceutically acceptable salt
- 25 thereof. Preferred prodrugs for use with CPG2 are described in the following US patents from Zeneca Limited and Cancer Research Campaign Technology Limited: US 5714148, US 5405990, 5587161 & 5660829.

In another aspect of the invention there is provided a method for the delivery of a cytotoxic drug to a site which comprises administering to a host a first component that comprises a gene construct as defined herein; followed by administration to the host of a second component that comprises a prodrug which can be converted into a cytotoxic drug by the heterologous enzyme encoded by the first component. A preferred method for delivery of

a cytotoxic drug to a site is one in which the first component comprises a gene encoding the heterologous enzyme CPG2; and the second component prodrug is selected from  $\underline{N}$ -(4- $[\underline{N},\underline{N}$ -bis(2-iodoethyl)amino]phenoxycarbonyl)- $\underline{L}$ -glutamic acid,  $\underline{N}$ -(4- $[\underline{N},\underline{N}$ -

 $bis (2-chloroethyl) a mino]-phenoxycarbonyl) - \underline{L}-glutamic-gamma-(3,5-dicarboxy) anilide or \underline{N}-$ 

5 (4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl)-<u>L</u>-glutamic acid or a pharmaceutically acceptable salt thereof.

## Abbreviations used herein include:

AAV Adeno-associated virus

ADEPT antibody directed enzyme prodrug therapy

AFP alpha-foetoprotein

AMIRACS Antimetabolite with Inactivation of Rescue Agents at

Cancer Sites

APS ammonium persulfate

b.p. base pair

BPB bromophenol blue

CDRs complementarity determining regions

CEA Carcinoma Embryonic Antigen

CL constant domain of antibody light chain

CPB carboxypeptidase B
CPG2 carboxypeptidase G2

CPG2 R6 carboxypeptidase G2 mutated to prevent

glycosylation on expression in eucaryotic cells, see

Example 1d

DAB substrate 3,3'-diaminobenzidine tetrahydrochloride

DEPC diethylpyrocarbonate

DMEM Dulbecco's modified Eagle's medium

ECACC European Collection of Animal Cell Cultures

EIA enzyme immunoassay

ELISA enzyme linked immunosorbent assay

FAS folinic acid supplemented

FCS foetal calf serum

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Fd

heavy chain of Fab, Fab' or F(ab')2 optionally

containing a hinge

**GDEPT** 

gene directed enzyme prodrug therapy

**HAMA** 

Human Anti Mouse Antibody

**HCPB** 

human carboxypeptidase B, preferably pancreatic

hinge (of an IgG)

a short proline rich peptide which contains the

cysteines that bridge the 2 heavy chains

HRPO or HRP

horse radish peroxidase

**IRES** 

internal ribosome entry site

MTX

methotrexate

**NCA** 

non-specific cross reacting antigen

**NCIMB** 

National Collections of Industrial and Marine

Bacteria

**OPD** 

ortho-phenylenediamine

PBS

phosphate buffered saline

PCR

polymerase chain reaction

**PGP** 

 $\underline{N}$ -(4- $[\underline{N},\underline{N}$ -bis(2-chloroethyl)amino]-

phenoxycarbonyl)-L-glutamic acid

preproCPB

proCPB with an N-terminal leader sequence

proCPB

CPB with its N-terminal pro domain

scFv

single chain Fv

SDS-PAGE

sodium dodecyl sulphate - polyacrylamide gel

electrophoresis

SSC

salt sodium citrate

TBS

Tris-buffered Saline

Temed

N,N,N',N'-tetramethylethylenediamine

TFA

trifluoroacetic acid

TRS

transcriptional regulatory sequence

**VDEPT** 

virus-directed enzyme prodrug therapy

VH

variable region of the heavy antibody chain

VK

variable region of the light antibody chain

In this specification conservative amino acid analogues of specific amino acid sequences are contemplated which retain the relevant biological properties of the component of the invention but differ in sequence by one or more conservative amino acid substitutions, deletions or additions. However the specifically listed amino acid sequences are preferred. Typical conservative amino acid substitutions are tabulated below.

Original	Exemplary Substitutions	Preferred Substitutions
Ala (A)	Val; Leu; Ile	Val
Arg (R)	Lys; Gln; Asn	Lys
Asn (N)	Gln; His; Lys; Arg	Gln
Asp (D)	Glu	Glu
Cys (C)	Ser	Ser
Gln (Q)	Asn	Asn
Glu (E)	Asp	Asp
Gly (G)	Pro	Pro
His (H)	Asn; Gln; Lys; Arg	Arg
Ile (I)	Leu; Val; Met; Ala; Phe;	Leu
	Norleucine	
Leu (L)	Norleucine; Ile; Val;	Ile
	Met; Ala; Phe	
Lys (K)	Arg; Gln; Asn	Arg
Met (M)	Leu; Phe; Ile	Leu
Phe (F)	Leu; Val; Ile; Ala	Leu
Pro (P)	Gly	Gly
Ser (S)	Thr	Thr
Thr (T)	Ser	Ser
Trp (W)	Тут	Tyr
Tyr (Y)	Trp; Phe; Thr; Ser	Phe

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Val (V)

Ile; Leu; Met; Phe;

Leu

Ala; Norleucine

Amino acid nomenclature is set out below.

Alanine	Ala	Α
Arginine	Arg	R
Asparagine	Asn	N
Aspartic Acid	Asp	D
Cysteine	Cys	С
Glutamic Acid	Glu	. Е
Glutamine	Gln	Q
Glycine	Gly	G
Histidine	His	Н
Isoleucine	Ile	I
Leucine	Leu	L
Lysine	Lys	K
Methionine	Met	· M
Phenylalanine	Phe	F
Proline	Pro	P
Serine	Ser	S
Threonine	Thr	Т
Tryptophan	Trp	W
Tyrosine	Tyr	Y
Valine	Val	V
Any Amino Acid	Xaa	X

5

In this specification nucleic acid variations (deletions, substitutions and additions) of specific nucleic acid sequences are contemplated which retain which the ability to hybridise under stringent conditions to the specific sequence in question. Stringent conditions are defined as 6xSSC, 0.1 % SDS at 60° for 5 minutes. However specifically listed nucleic acid

sequences are preferred. It is contemplated that chemical analogues of natural nucleic acid structures such as "peptide nucleic acid" (PNA) may be an acceptable equivalent, particularly for purposes that do not require translation into protein (Wittung (1994) Nature 368, 561).

The invention will now be illustrated by reference to the following non-limiting 5 Examples. Temperatures are in degrees Celsius.

Figure 1 shows a representation of the fusion gene construct comprising A5B7 antibody heavy chain Fd fragment linked at its C-terminus via a flexible (G4S)3 peptide linker to the N-terminus of CPG2 polypeptide. SS represents the signal sequence. L represents a linker sequence. CPG2/R6 represents CPG2 with its glycosylation sites nullified through mutation as explained in the text.

Figure 2a shows a representation of (Fab-CPG2)<sub>2</sub> fusion protein with dimerisation taking place through non-covalent bonding between two CPG2 molecules.

Figure 2b shows a representation of a F(ab')2 antibody fragment.

Figure 3 shows a cell based ELISA assay of secreted fusion protein material. Only the CEA positive line has increased levels of binding with increasing amounts of added fusion protein whereas the CEA negative cell line has only constant background binding levels throughout. The vertical axis represents optical density readings measured at 490 nm and the horizontal axis the amount of added fusion protein measured in ng of protein. The graph shows data obtained from an experiment where a number of cell lines and a negative control (no cells) were incubated with increasing amounts of fusion protein using the cell assay described in Example 6. The results show that only the LoVo (CEA positive) cell line showed an increasing OD490 reading corresponding to increasing amounts of addes fusion protein. All other cell lines (CEA negative) and the control (no cells) showed only a background OD490 nm reading which did not increase with the addition of fusion protein. These results provide evidence that the fusion protein material binds specifically to a CEA positive cell line in a dose dependant manner and do not bind to CEA negative lines.

Figure 4 shows retention of secreted fusion protein to recombinant LoVo tumour cells. The vertical axis represents optical density readings measured at 490 nm and the horizontal axis the amount of added anti-CEA antibody (IIE6) measured in ng/ml of protein. The experiment 30 was performed as described in Example 7 using three different cell lines, recombinant LoVo and Colo320DM lines (which themselves secrete fusion protein) and a contol parental LoVo line which does not secrete fusion protein. Firstly, the cell lines were fixed, and washed to

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remove the existing supernatant and any unbound material after which increasing concentrations of the anti-CEA antibody (IIE6) were added to the fixed cells. The assay was developed as described in the text to determine the level of retention of any secreted material and whether further added antibody would increase the signal. The results showed that 5 whithout added anti-CEA antibody the control parental Lovo line exhibited only a backgroundOD490 nm reading (as expected) whereas the recombinant LoVo line gave a very strong OD 490 nm reading indicating that the fusion protein material was being retained on the CEA positive LoVo cells. The CEA negative recombinant Colo320DM gave a much weaker reading than the LoVo cells but the signal was higher than background (possibly due 10 to none fixing of the secreted antibody early in the assay method). Increasing concentrations of the anti-CEA antibody (IIE6) added to the fixed cells showed a dose related response in the case of the parental LoVo cells thus indicating that they are CEA positive and can bind CEA binding material (such as the fusion protein if present or added). The recombinant Colo320DM and LoVo cells showed little increase in overall OD490 signal with increasing 15 amounts of added antibody with the exception of the LoVo cells which appear to show a slight response at the highest antibody dose. Since the recombinant Colo320DM are CEA negative no increase in signal due to anti-CEA antibody the results for these cells would be expected. In the case of the recombinant LoVo cells the addition signal due the amounts of antibody added in this assay may be swamped except at the highest dose due to the relative 20 strength of the original signal.

Figure 5 shows retention of secreted fusion protein to recombinant LoVo tumour cells. The vertical axis represents median tumour volume (cm³) and the horizontal axis time in day after dosing of the prodrug. The experiment was performed as described in Example 12 using 60 mg/kg doses of prodrug. The results show that the control GAD(c) (none prodrug treated) tumours grew to 6 times their initial size by 11 days (post-dose day) at which time the tumours were harvested. The prodrug treated tumours GAD(d) show a significantly slower growth rate and by day 16 (post-dose day) have only reached 3 times their initial size. This

In the Examples below, unless otherwise stated, the following methodology and 30 materials have been applied.

data indicates at least an 11 day tumour growth delay.

DNA is recovered and purified by use of GENECLEAN™ II kit (Stratech Scientific Ltd. or Bio 101 Inc.). The kit contains: 1) 6M sodium iodide; 2) a concentrated solution of

sodium chloride, Tris and EDTA for making a sodium chloride/ethanol/water wash; 3) Glassmilk- a 1.5 ml vial containing 1.25 ml of a suspension of a specially formulated silica matrix in water. This is a technique for DNA purification based on the method of Vogelstein and Gillespie published in Proceedings of the National Academy of Sciences USA (1979) Vol

- 5 76, p 615. Briefly, the kit procedure is as follows. To 1 volume of gel slice is added 3 volumes of sodium iodide solution from the kit. The agarose is melted by heating the mix at 55° for 10 min then Glassmilk (5-10 ml) is added, mixed well and left to stand for 10 min at ambient temperature. The glassmilk is spun down and washed 3 times with NEW WASHTM (0.5 ml) from the kit. The wash buffer is removed from the Glassmilk and DNA is eluted by
- 10 incubating the Glassmilk with water (5-10 ml) at 55° for 5-10 min. The aqueous supernatant containing the eluted DNA is recovered by centrifugation. The elution step can be repeated and supernatants pooled.

Competent E. coli DH5α cells were obtained from Life Technologies Ltd (MAX<sup>TM</sup> efficiency DH5α competent cells).

- Mini-preparations of double stranded plasmid DNA were made using the RPMTM DNA preparation kit from Bio101 Inc. (cat. No 2070-400) or a similar product the kit contains alkaline lysis solution to liberate plasmid DNA from bacterial cells and glassmilk in a spinfilter to adsorb liberated DNA which is then eluted with sterile water or 10mM Tris-HCl, 1mM EDTA, pH 7.5.
- The standard PCR reaction contains 100 ng of plasmid DNA (except where stated), 5 μl dNTPs (2.5 mM), 5 μl 10x Enzyme buffer (500 mM KCl, 100 mM Tris pH 8.3), 15mM MgCl<sub>2</sub> and 0.1 % gelatin), 1 μl of a 25 pM/ μl stock solution of each primer, 0.5 μl thermostable DNA polymerase and water to obtain a volume of 50 μl. Standard PCR conditions were: 15 cycles of PCR at 94° for 90 s; 55° for 60 s; 72° for 120 s, ending the last 25 cycle with a further 72° for 10 min incubation.

AMPLITAQ™, available from Perkin-Elmer Cetus, is used as the source of thermostable DNA polymerase.

General molecular biology procedures can be followed from any of the methods described in "Molecular Cloning - A Laboratory Manual" Second Edition, Sambrook, Fritsch and Maniatis (Cold Spring Harbor Laboratory, 1989).

Serum free medium is OPTIMEM™ I Reduced Serum Medium, GibcoBRL Cat. No.

31985. This is a modification of Eagle's Minimum Essential Medium buffered with Hepes and sodium bicarbonate, supplemented with hypoxanthine, thymidine, sodium pyruvate, <u>L</u>glutamine, trace elements and growth factors.

LIPOFECTIN<sup>TM</sup> Reagent (GibcoBRL Cat. No. 18292-011) is a 1:1 (w/w) liposome
5 formulation of the cationic lipid N-[1-(2,3-dioleyloxy)propyl]-n,n,n-trimethylammonium
chloride (DOTMA) and dioleoyl phosphatidylethanolamine (DOPE) in membrane filtered
water. It binds spontaneously with DNA to form a lipid-DNA complex - see Felgner et al. in
Proc. Natl. Acad. Sci. USA (1987) 84, 7431.

G418 (sulphate) is GENETICIN<sup>TM</sup>, GibcoBRL Cat. No 11811, an aminoglycoside antibiotic related to gentamicin used as a selecting agent in molecular genetic experiments;

For the CEA ELISA each well of a 96 well immunoplate (NUNC MAXISORB<sup>TM</sup>) was coated with 50ng CEA in 50 mM carbonate/bicarbonate coating buffer pH9.6 (buffer capsules - Sigma C3041) and incubated at 4° overnight. The plate was washed three times with PBS-TWEEN<sup>TM</sup> (PBS + 0.05 % TWEEN<sup>TM</sup> 20) and then blocked 150 μl per well of 1 % BSA in PBS-TWEEN<sup>TM</sup> for 1 hour at room temperature. The plate was washed three times with PBS-TWEEN<sup>TM</sup>, 100 μl of test sample added per well and incubated at room temperature for 2 hours. The plate was washed three times with PBS-TWEEN<sup>TM</sup>, 100 μl per well of a 1/500 dilution of HRPO-labelled goat anti-human kappa antibody (Sigma A 7164) was added in 1 % BSA in PBS-TWEEN<sup>TM</sup> and incubated at room temperature on a rocking platform for at least 1 hour. The plate was washed three times with PBS-TWEEN<sup>TM</sup> and then once more with PBS. To detect binding, add 100μl per well of developing solution (one capsule of phosphate-citrate buffer - Sigma P4922 - dissolved in 100 ml H<sub>2</sub>O to which is added one 30 mg tablet *o*-phenylenediamine dihydrochloride - Sigma P8412) and incubated for up to 15 minutes. The reaction was stopped by adding 75 μl 2M H<sub>2</sub>SO<sub>4</sub>, and absorbance read at 490 nm.

The CEA ELISA using an anti CPG2 reporter antibody was essentially as above but instead of HRPO-labelled goat anti-human kappa antibody an 1/1000 dil. of a rabbit anti-CPG2 polyclonal sera was added, in 1 % BSA in PBS-TWEEN<sup>TM</sup> and incubated at room temperature on a rocking platform for at 2 hours. The plate was washed three times with 30 PBS-TWEEN<sup>TM</sup>. A 1/2000 dilution of a goat anti-rabbit HRPO labelled antibody (Sigma A-6154) was then added and incubated at room temperature on a rocking platform for 1 hour, the

plate was washed three times with PBS-TWEEN™ and once with PBS. To detect binding add 100µl per well developing solution (one capsule of phosphate-citrate buffer - Sigma P4922 - dissolved in 100 ml H<sub>2</sub>O to which is added one 30 mg tablet *o*-phenylenediamine dihydrochloride - Sigma P8412) and incubated for up to 15 minutes. The reaction was 5 stopped by adding 75µl 2M H<sub>2</sub>SO<sub>4</sub>, and absorbance read at 490nm.

Western blot analysis of transfection supernatants was performed as follows.

10 % mini gels for analysis of fusion protein transfections were prepared using a suitable mini gel system (HOEFER MIGHTY SMALL<sup>TM</sup>). 10 % running gel is: 20 ml acrylamide, 6 ml 10 x running gel buffer; 34 ml H<sub>2</sub>O; 300 ml 20 % SDS; 600 μl APS; 30 μl Temed.

10 Running gel buffer 10x is 3.75 M Tris pH 8.6. 6 % stacking gel is: 9 ml acrylamide; 4.5 ml 10x stacking gel buffer; 31.5 ml H<sub>2</sub>O; 225 μl 20 % SDS 450 μl 10 % APS; 24 μl Temed). Stacking gel buffer 10x is 1.25 M Tris pH 6.8. Electrophoresis buffer 5x for SDS/PAGE is 249 mM Tris, 799 mM glycine, 0.6 % w/v SDS (pH not adjusted).

Preparation of samples 2 x Laemmli buffer is 0.125 M Tris; 4 % SDS; 30 %

15 glycerol; 4 M urea; 0.002 % BPB optionally containing 5 % β-mercaptoethanol.

Supernatants: 25 μl sample + 25 μl 2 x Laemmli buffer; 40 μl loaded. Standards F(ab')<sub>2</sub>

and CPG2: 2 μl of 10 ng/ml of standard; 8 μl of H<sub>2</sub>O; 10 μl 2x Laemmli buffer (mercaptoethanol); 20 μl loaded. Molecular weight markers (Amersham RAINBOWTM): 8

μl sample; 8 μl 2x Laemmli buffer (+ mercaptoethanol): 16 μl loaded. Running conditions:

20 30 milliamps until dye front at bottom of gel(approx. 1 hour). Blotting: using a semi dry blotter (LKB) onto nitrocellulose membrane. Milliamps = 0.7 x cm<sup>2</sup>, for 45 minutes.

Blocking: 5 % dried skimmed milk in PBS-TWEENTM for 40 minutes.

Detection of F(ab')<sub>2</sub>: goat anti human kappa light chain HRPO labelled antibody, 1/2500 in 0.5 % dried skimmed milk in PBS-TWEEN™ incubated overnight.

Detection of CPG2: mouse anti-CPG2 monoclonal (1/2000 in 0.5 % dried skimmed milk in PBS-TWEEN<sup>TM</sup> incubated overnight; goat anti mouse kappa light chain HRPO labelled antibody -Sigma 674301- (1/10000 in 0.5 % dried skimmed milk in PBS-TWEEN<sup>TM</sup>) incubated for at least 2 hours.

Development of Blot: Chemiluminescence detection of HRPO based on luminol
30 substrate in the presence of enhancer was used (Pierce SUPERSIGNAL™ Substrate).
Substrate working solution was prepared as follows: recommended volume: 0.125 ml/cm² of

blot surface. Mix equal volumes of luminol/enhancer solution and stable peroxide solution, incubate blot with working solution for 5-10 minutes, remove solution and place blot in a membrane protector and expose against autoradiographic film (usually between 30 seconds and 5 minutes).

- Microorganism deposits: Plasmid pNG3-Vkss-HuCk was deposited at The National Collections of Industrial and Marine Bacteria (NCIMB), 23 St Machar Drive, Aberdeen AB2 1RY, Scotland, United Kingdom on 11-April-1996 under deposit reference number NCIMB 40798 in accordance with the Budapest Treaty. Plasmid pNG4-VHss-HuIgG2CH1' was deposited at The National Collections of Industrial and Marine Bacteria (NCIMB), 23 St
- 10 Machar Drive, Aberdeen AB2 1RY, Scotland, United Kingdom on 11-April-1996 under deposit reference number NCIMB 40797 in accordance with the Budapest Treaty. Plasmid pNG3-Vkss-HuCk-NEO was deposited at The National Collections of Industrial and Marine Bacteria (NCIMB), 23 St Machar Drive, Aberdeen AB2 1RY, Scotland, United Kingdom on 11-April-1996 under deposit reference number NCIMB 40799 in accordance with the
- Budapest Treaty. Plasmid pICI266 was deposited under accession number NCIMB 40589 on 11Oct93 under the Budapest Treaty at the National Collections of Industrial and Marine Bacteria Limited (NCIMB), 23 St. Machar Drive, Aberdeen, AB2 1RY, Scotland, U.K.

Typsinisation: Trypsin EDTA (Gibco BRL 45300-019) and Hanks balanced salt solution (HBSS; Gibco BRL 14170-088) were pre-warmed in a 37° waterbath. Existing 20 media was removed from cultures and replaced with a volume of HBSS (which is half the previous media volume) and the layer of cells washed by carefully rocking the plate or flask so as to remove any residual serum containing media. The HBSS was removed and a volume of Trypsin solution (which is one quarter of the original media volume) added, with gently rocking the flask to ensure the cell layer was completely covered and left for 5 min. Trypsin 25 was inactivated by addition of of the appropriate normal culture media (2x the volume of the trypsin solution). The cell suspension was then either cell counted or further diluted for continued culture depending on the procedure to be performed.

Heat Inactivation of Foetal Calf Serum (FCS): FCS (Viralex A15-651 accredited batch - Non European) was stored at -20°. For use, the serum was completely thawed at 4° overnight. The next day, the serum was incubated for 15 min in a 37° waterbath and then transferred to a 56° waterbath for 15 min. The serum was removed and allowed to cool to room temperature before it was split in to 50 ml aliquots and stored at -20°C

Normal DMEM Media (using Gibco BRL components): To 500 ml DMEM (41966-086) add 12.5ml Hepes (15630-056); 5ml NEAA (11140-035); 5 ml pen/strep (10378-016); and 50 ml heat inactivated FCS.

FAS Media (using Gibco BRL components unless stated otherwise): 490 ml DMEM 5 (41966-086); 12.5 ml Hepes (15630-056); 5ml non-essential amino acids (11140-035); 5 ml pen/strep (10378-016); 5 ml vitamins (11120-037); 5ml basal amino acids (51051-019); Folinic Acid (Sigma F8259) to a final media concentation of 10 μg/ml; 50 ml heat inactivated FCS; 5 ml dNTP mix; and G418 50 mg/ml stock solution (to produce the appropriate selection concentration).

dNTP mix: 35mg G (Sigma G6264), 35mg C (Sigma C4654), 35mg A (Sigma A4036), 35mg U (SigmaU3003), 125mg T (Sigma T1895) were dissolved in 100ml water, filter sterilised, and stored at -20°.

G418 Selection: for LoVo cells (ATCC CCL 229) selection was performed at 1.25 mg/ml, for HCT116 (ATCC CCL 247) cells and for Colo320DM (ATCC CCL 220) cells selection was performed at 1.5 mg/ml unless stated otherwise.

BLUESCRIPT<sup>TM</sup> vectors were obtained from Stratagene Cloning Systems.

Tet-On gene expression vectors were obtained from Clontech (Palo Alto, California) cat. no. K1621-1.

Unless stated otherwise or apparent from the context used, antibody-CPG2 fusion 20 constructs referred to in the Examples use mutated CPG2 to prevent glycosylation.

#### Example 1

## Construction of an (A5B7 Fab-CPG2)2 fusion protein

The construction of a (A5B7 Fab-CPG2)<sub>2</sub> enzyme fusion was planned with the aim of obtaining a bivalent human carcinoembryonic antigen (CEA) binding molecule which also exhibits CPG2 enzyme activity. To this end the initial construct was designed to contain an A5B7 antibody heavy chain Fd fragment linked at its C-terminus via a flexible (G4S)<sub>3</sub> peptide linker to the N-terminus of the CPG2 polypeptide (Figure 1).

The antibody A5B7 binds to human carcinoembryonic antigen (CEA) and is

30 particularly suitable for targeting colorectal carcinoma or other CEA antigen bearing cells (the importance of CEA as a cancer associated antigen is reviewed by Shively, J.E. and Beatty,

J.D. in "CRC Critical Reviews in Oncology/Hematology", vol 2, p355-399, 1994).

The CPG2 enzyme is naturally dimeric in nature, consisting of two associated identical polypeptide subunits. Each subunit of this molecular dimer consists of a larger catalytic domain and a second smaller domain that forms the dimer interface.

In general, antibody (or antibody fragment)-enzyme conjugate or fusion proteins

5 should be at least divalent, that is to say capable of binding at least 2 tumour associated antigens (which may be the same or different). In the case of the (A5B7 Fab-CPG2)<sub>2</sub> fusion protein, dimerisation of the enzyme component takes place after expression, as with the native enzyme, thus forming an enzymatic molecule which contains two Fab antibody fragments (and is thus bivalent with respect to antibody binding sites) and two molecules of CPG2

10 (Figure 2a).

#### a) Cloning of the A5B7 antibody genes

Methods for the preparation, purification and characterisation of recombinant murine A5B7 F(ab')<sub>2</sub> antibody have been published (International Patent Application, Zeneca Limited, WO 96/20011, see Reference Example 5 therein). In Reference Example 5, section f thereof, the A5B7 antibody genes were cloned into vectors of the GS-SYSTEM<sup>TM</sup> (Celltech), see International Patent Applications WO 87/04462, WO 89/01036, WO 86/05807 and WO 89/10404, with the A5B7 Fd cloned into pEE6 and the light chain into pEE12. These vectors were the source of the A5B7 antibody genes for the construction of the A5B7 Fab-CPG2 fusion protein.

#### 20 b) Chimaeric A5B7 vector constructs

The A5B7 murine antibody variable regions were amplified by PCR from the pEE6 and pEE12 plasmid vectors using appropriate PCR primers which included the necessary restriction sites for direct in frame cloning of the heavy and light chain variable regions into the vectors pNG4-VHss-HulgG2CH1' (NCIMB deposit no. 40797) and pNG3-Vkss-HuCk-

25 NEO (NCIMB deposit no. 40799) respectively. The resulting vectors were designated pNG4/A5B7VH-IgG2CH1' (A5B7 chimaeric heavy chain Fd') and pNG3/A5B7VK-HuCK-NEO (A5B7 chimaeric light chain).

#### c) Cloning of the CPG2 gene

The CPG2 coding gene may be obtained from Centre for Applied Microbiology and Research, Porton Down, Salisbury, Wiltshire SP4 0JG, United Kingdom. CPG2 may also be obtained by recombinant techniques. The nucleotide coding sequence for CPG2 has been published by Minton, N.P. et al., Gene, (1984) 31, 31-38. Expression of the coding sequence

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has been reported in *E.coli* (Chambers, S.P. et al., Appl. Microbiol, Biotechnol. (1988), 29, 572-578) and in *Saccharomyces cerevisiae* (Clarke, L. E. et al., J. Gen Microbiol, (1985) 131, 897-904). In addition the CPG2 gene may be produced as a synthetic DNA construct by a variety of methods and used as a source for further experiments. Total gene synthesis has 5 been described by M. Edwards in Am. Biotech. Lab (1987), 5, 38-44, Jayaraman et al. (1991) Proc. Natl. Acad. Sci. USA 88, 4084-4088, Foguet and Lubbert (1992) Biotechniques 13, 674-675 and Pierce (1994) Biotechniques 16, 708.

In preparation for the cloning the CPG2 gene the vector pNG3-Vkss was constructed which is a simple derivative of pNG3-Vkss-HuCk-NEO (NCIMB deposit no. 40799). This vector was constructed by first removing the Neomycin gene (since it contained an EcoRI restriction enzyme site) by digestion with the restriction enzyme XbaI, after which the vector fragment was isolated and then religated to form the plasmid pNG3/Vkss-HuCk. This intermediate vector was digested with the enzymes SacII and EcoRI, which excised the HuCk gene fragment. The digest was then loaded on a 1 % agarose gel and the excised fragment separated from the remaining vector after which the vector DNA was cut from the gel and purified. Two oligonucleotides CME 00261 and CME 00262 (SEQ ID NO: 1 and 2) were designed and synthesised. These two oligonucleotides were hybridised by adding 200 pmoles of each oligonucleotide into a total of 30 µl of H<sub>2</sub>O, heating to 95° and allowing the solution to cool slowly to 30°. 100 pmoles of the annealed DNA product was then ligated directly into 20 the previously prepared vector and the ligation mix transformed into *E. coli*. In the clones obtained, the introduction of the DNA "cassette" produced a new polylinker sequence in preparation for the subsequent CPG2 gene cloning to produce the vector pNG3-Vkss.

The CPG2 structural gene encoding amino acid residues Q26-K415 inclusive was amplified by PCR using appropriate DNA oligonucleotide primers and standard PCR reaction conditions. The reaction product was analysed using a 1 % agarose gel, a band of the expected size (approximately 12000 b.p.) was excised, purified and eluted in 20µl H<sub>2</sub>O. This material was then digested using the restriction enzyme SacII, after which the reaction was loaded on a 1 % agarose gel and a band of the expected size (approximately 250 b.p.) was excised and subsequently purified. This fragment was ligated into the plasmid vector pNG3VKss, which had been previously digested with the restriction enzyme SacII, dephosphorylated, run on a 1 % agarose gel, the linearised vector band excised, purified, and the ligation mix transformed

into *E.coli*. The resultant clones were analysed for the presence and orientation of the CPG2 SacII fragment by DNA restriction analysis using the enzymes BgIII and FseI. Clones which appeared to have a fragment of the correct size and orientation were confirmed by DNA sequencing. This intermediate plasmid was called pNG3-Vkss-SacIICPG2frag. This

5 plasmid was digested with the restriction enzymes by AgeI and EcoRI, dephosphorylated and the vector fragment isolated. The original CPG2 gene PCR product was also digested with AgeI and EcoRI, an approximately 1000 bp. fragment isolated, ligated and transformed into *E.coli*. The resulting clones were analysed for a full length CPG2 gene (approximately 1200 bp.) by digestion with the restriction enzymes HindIII and EcoRI; clones with the correct size insert were sequenced to confirm identity. Finally, this plasmid (pNG3/Vkss-CPG2) was digested with XbaI, dephosphorylated, a vector fragment isolated and the XbaI Neomycin gene fragment (approximately 1000 bp. which had also been isolated in the earlier stages) religated into the plasmid and transformed into *E.coli*. Resulting clones were checked for the presence and orientation of the Neomycin gene by individual digests with the enzymes XbaI and EcoRI. This vector was called pNG3-Vkss-CPG2-NEO.

## d) Construction of the CPG2 R6 variant

The plasmid pNG3-Vkss/CPG2-NEO was used as a template for the PCR mutagenesis of the CPG2 gene in order to mutate 3 potential glycosylation sites which had been identified within the natural bacterial enzyme sequence. The putative amino acid glycosylation sites (N-

- 20 X-T/S) were observed at positions 222 (N-I-T), 264 (N-W-T), and 272 (N-V-S) using the positional numbering published by Minton, N.P. et al., in Gene, (1984) 31, 31-38. The asparagine residue (N) of the 3 glycosylation sites was mutated to glutamine (Q) thus negating the glycosylation sites to avoid any glycosylation events affecting CPG2 expression or enzyme activity.
- A PCR mutagenesis technique in which all 3 sites were mutated in a single reaction series was used to create the CPG2 R6 gene variant. The vector pNG3/Vkss/CPG2-NEO was used as the template for three initial PCR reactions. Reaction R1 used synthetic oligonucleotide sequence primers CME 00395 and CME 00397 (SEQ ID NOS: 3 and 4), reaction R2 used synthetic oligonucleotide sequence primers CME 00395 and CME 00399 30 (SEQ ID NOS: 3 and 5) and reaction R3 used synthetic oligonucleotide sequence primers
- 30 (SEQ ID NOS: 3 and 5) and reaction R3 used synthetic oligonucleotide sequence primers CME 00396 and CME 00400 (SEQ ID NOS: 6 and 7). The products of PCR reactions R1 and R2 contained the mutated 222 and 264 + 272 glycosylation sites respectively, with the R3

product being a copy of the C-terminal segment of the CPG2 gene. The R2 and R3 products (R2 approximately 750 bp; R3 approximately 360 bp), after agarose gel separation and purification, were joined in a further PCR reaction. Mixtures of varying amounts of the products R2 and R3 were made and PCR reactions performed using the synthetic oligonucleotides CME 00395 and CME 00396 (SEQ ID NOS: 3 and 6). The resulting product R4 (approximately 1200bps) was again PCR amplified using the oligonucleotides CME 00398 and CME 00396 (SEQ ID NOS: 8 and 6). The resulting product R5 (approximately 600 bp.) was joined to product R1 (approximately 620 b.p.) in a final PCR reaction performed using the oligonucleotides CME 00395 and CME 00396 (SEQ ID NOS: 3 and 6). The resulting PCR product R6 (approximately 1200 bp), which now contained all three mutated glycosylation sites, could be cloned (after digestion with the restriction

three mutated glycosylation sites, could be cloned (after digestion with the restriction enzymes AgeI and BsrGI and isolation of the resultant fragment) into the vector pNG3/Vkss-CPG2-Neo.(which had been previously cut with the restriction enzymes AgeI and Bsr GI and subsequently isolated). This created the desired DNA (SEQ ID NO: 9) encoding CPG2/R6 protein sequence (SEQ ID NO: 10) within the expression vector pNG3/Vkss-CPG2 R6-NEO.

# e) Construction of the A5B7 heavy chain Fd-CPG2 fusion protein gene

The heavy chain antibody fragment and the CPG2 enzyme genes were both obtained by PCR amplification of plasmid templates. The plasmid pNG4/A5B7VH-IgG2CH1' was amplified with primers CME 00966 (SEQ ID NO: 11) and CME 00969 (SEQ ID NO: 12) to obtain the A5B7 Fd component (approximately 300 b.p.) and the plasmid pNG3/Vkss/CPG2 R6-NEO was amplified with primers CME 00967 (SEQ ID NO: 13) and CME 00968 (SEQ ID NO: 14) to obtain the enzyme component (approximately 1350 b.p.). In each case the PCR reaction product was loaded and separated on a 1 % agarose gel, a band of the correct product size excised, subsequently purified and eluted in 20 µl H<sub>2</sub>O.

A further PCR reaction was performed to join (or splice) the two purified PCR reaction products together. Standard PCR reaction conditions were used with varying amounts (between 0.5 to 2 μl) of each PCR product but utilising 25 cycles (instead of the usual 15 cycles). The reaction product was analysed using a 1 % agarose gel and a band of the expected size (approximately 1650 b.p.) was excised, purified and eluted in 20 μl H<sub>2</sub>O. This material was then digested using restriction enzymes Nhel and BamHI, after which a

band of the expected size (approximately 1600 b.p.) was recovered and purified. The vector pNG4/A5B7VH-IgG2CH1' was prepared to receive the above PCR product by digestion with restriction enzymes NheI and BamHI, after which the DNA was dephosphorylated and the larger vector band was separated from the smaller NheI/Bam HI fragment. The vector band was recovered, purified and subsequently the similarly restricted PCR product was ligated in to the prepared vector and the ligation mix transformed into *E. coli*. DNA was prepared from the clones obtained and subsequently sequenced to confirm the fusion gene sequence. A number of the clones were found to be correct and one of these clones (designated R2.8) was re-named pNG4/A5B7VH-IgG2CH1/CPG2 R6 (SEQ ID NO: 15 and SEQ ID NO: 16).

## 10 f) Co-transfection, transient expression

The plasmids pNG4/A5B7VH-IgG2CH1/CPG2 R6 (encoding the antibody chimaeric Fd-CPG2 fusion protein) and pNG3/A5B7VK-HuCK-NEO (encoding the antibody chimaeric light chain; SEQ ID NO: 17 and SEQ ID NO: 18) were co-transfected into COS-7 cells using a LIPOFECTIN<sup>TM</sup> based procedure as described below. COS7 cells are seeded into a 6 well

- plate at 2x105cells/2 ml/well, from a subconfluent culture and incubated overnight at 370, 5 % CO2. A LIPOFECTINTM/ serum free medium mix is made up as follows: 12 ml LIPOFECTINTM plus 200 ml serum free medium and incubated at room temperature for 30 minutes. A DNA/serum free medium mix is made up as follows: 4 mg DNA (2 mg of each construct) plus 200 ml serum free medium. 200 ml of the LIPOFECTINTM/ serum free
- 20 medium mix is then added to the DNA mix and incubated for 15 minutes room temperature. 600 ml of serum free medium was then added to each sample. The cells were washed once with 2 ml serum free medium and then the 1 ml LIPOFECTIN™/DNA mix is added to the cells and incubated for 5 hours, 37°, 5 % CO₂. The LIPOFECTIN™/DNA mix was removed from the cells and normal growth media added after which the cells were incubated for 72 hours, 37°, 5 % CO₂. The cell supernatants were harvested.

## g) Analysis of Antibody-Enzyme Fusion Protein

The supernatant material was analysed for the presence of antibody fusion protein using a CEA-binding ELISA using an anti human kappa light chain reporter antibody (for presence of antibody), a CEA-binding ELISA using an anti-CPG2 reporter antibody (for

30 presence of CEA bound CPG2 fusion protein), a HPLC based CPG2 enzyme activity assay (to measure specific CPG2 activity) and SDS/PAGE followed by Western blotting (using either anti human kappa light chain reporter or anti CPG2 reporter antibodies) to detect expressed material.

The HPLC based enzyme activity assay clearly showed CPG2 enzyme activity to be present in the cell supernatant and both the anti-CEA ELISA assays exhibited binding of protein at levels commensurate with a bivalent A5B7 antibody molecule. The fact that the anti-CEA ELISA detected with an anti-CPG2 reporter antibody also exhibited clear CEA binding indicated that not only antibody but also antibody-CPG2 fusion protein was binding CEA.

Western blot analysis with both reporter antibody assays clearly displayed a fusion protein subunit of the expected approximately 90 kDa size with no degradation or smaller 10 products (such as Fab or enzyme) observable.

Since CPG2 is known only to exhibit enzyme activity when it is in a dimeric state and since only antibody enzyme fusion protein is present, this indicates that the 90 kDa fusion protein (seen under SDS/PAGE conditions) dimerises via the natural CPG2 dimerisation mechanism to form a 180 kDa dimeric antibody-enzyme fusion protein molecule (Figure 2a) in "native" buffer conditions. Furthermore, this molecule exhibits both CPG2 enzymatic activity and CEA antigen binding properties which do not appear to be significantly different in the fusion protein compared with enzyme or antibody alone.

- h) Use of expressed fusion protein and CPG2 prodrug in an *in vitro* cytotoxicity assay
- An in vitro cell killing assay was performed in which the (A5B7-CPG2 R6)<sub>2</sub> fusion protein was compared to a "conventional" A5B7 F(ab')<sub>2</sub>-CPG2 conjugate formed through linking A5B7 F(ab')<sub>2</sub> to CPG2 with a chemical heterobifunctional reagent. In each case material displaying equal amounts of CPG2 enzyme activity or equal amounts of antibody-CPG2 protein were incubated with LoVo, CEA bearing, tumour cells. The cells were then washed to remove unbound protein material and subsequently resuspended in medium containing a CPG2 phenol prodrug (PGP, see Example 2 below) for a period of 1 hr, after which the cells were washed, resuspended in fresh media and left to proliferate for 4 days. Finally the cells were treated with SRB stain and their numbers determined.

The results obtained clearly showed that the (A5B7-CPG2 R6)<sub>2</sub> fusion protein

30 (together with prodrug) caused at least equivalent cell kill and resulted in lower numbers of cells at the end of the assay period than the equivalent levels of A5B7 F(ab)<sub>2</sub>-CPG2 conjugate (with the same prodrug). Cell killing (above basal control levels) can only occur if the prodrug

is converted to active drug by the CPG2 enzyme (and since the cells are washed to remove unbound protein, only cell bound enzyme will remain at the stage where the prodrug is added). Thus this experiment shows that at least as much of the A5B7-CPG2 R6 fusion protein remains bound compared with conventional A5B7 F(ab)<sub>2</sub>-CPG2 conjugate as a greater 5 degree of cell killing (presumably due to higher prodrug to drug conversion) occurs.

i) Construction of a coexpression fusion protein vector for use in transient and stable cell line expression

For a simpler transfection methodology and the direct coupling of both expression cassettes to a single selection marker, a co-expression vector for fusion protein expression was 10 constructed using the existing vectors pNG4/A5B7VH-IgG2CH1/CPG2 R6 (encoding the antibody Fd-CPG2 fusion protein) and pNG3/A5B7VK-HuCK-NEO (encoding the antibody light chain). The pNG4/A5B7VH-IgG2CH1/CPG2 R6 plasmid was first digested with the restriction enzyme Scal, the reaction loaded on a 1 % agarose gel and the linear vector band excised from the gel and purified. This vector DNA was then digested with restriction 15 enzymes BglII and BamHI, the reaction loaded on a 1 % agarose gel, the desired band (approximately 2700 bp) recovered and purified. The plasmid pNG3/A5B7VK-HuCK-NEO was digested with the restriction enzyme BamHI after which the DNA was dephosphorylated then subsequently loaded on a 1 % agarose gel and the vector band excised from the gel and purified. The heavy chain expression cassette fragment was ligated in to the prepared vector 20 and the ligation mix transformed into E. coli. The orientation was checked by a variety of restriction digests and clones selected which had the heavy chain cassette in the same direction as that of the light chain. These plasmids were termed pNG3-A5B7-CPG2/R6coexp.-NEO.

## j) Gene switches for protein expression

It is foreseen that *in vitro* expression of CPG2 and CPG2 fusion proteins in mammalian cells may degrade media folates leading to slow cell growth or cell death. The high activity of the CPG2 enzyme is likely to make such a folate deficiency difficult to overcome by media supplementation. However, it is thought that in the case of CPG2 or CPG2 fusion protein expression from mammalian cells *in vivo*, it is unlikely that such problems will occur, since the cells would be constantly replenished with all growth requirements by the normal circulatory and cellular mechanisms.

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A number of options to avoid possible *in vitro* folic acid depletion problems have been considered. One of these solutions involve the use of tightly controlled but inducible gene switch systems such as the "TET on" or "TET off" switches (Grossen, M. et al (1995) Science 268: 1766-1769) or the ecdysone/ muristerone A switch (No, D. et al (1996) PNAS 93:3346-3351). Such systems enable precisely controlled expression of a gene of interest and allow stable transformation of mammalian cells with genes encoding toxic or potentially deleterious expression products. A gene switch would allow recombinant stable cell lines incorporating CPG2 fusion genes to be potentially more easily established, maintained and expanded for protein expression and seeding cultures for *in vivo* tumour growth studies.

10

## Example 2

HCT116 tumour cells expressing the antibody-enzyme fusion protein are selectively killed in vitro by a prodrug.

HCT116 colorectal tumour cells (ATCC CCL 247) transfected with the antibody15 CPG2 fusion protein gene of Example 1 can be selectively killed by a prodrug that is
converted by the enzyme into an active drug.

To demonstrate this, control non-transfected HCT116 cells or HCT116 cells transfected with the antibody-CPG2 fusion protein gene, are incubated with either the prodrug, 4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl-L-glutamic acid (PGP; Blakey et al, Br. J. cancer 72, 1083, 1995) or the corresponding drug released by CPG2, 4-[N,N-bis(2-chloroethyl)amino] phenol. PGP prodrug and drug over the concentration range of 5 X 10<sup>-4</sup> to 5 X 10<sup>-8</sup> M are added to 96 well microtitre plates containing 1000-2,500 HCT116 cells/well, for 1 hr at 37°. The cells are then washed and incubated for a further three days at 37°. After washing to remove dead cells, TCA is then added and the amount of cellular protein adhering to the plates is assessed by addition of SRB dye as described by Skehan et al ( J. Natl. Cancer Inst. 82, 1107, 1990). Potency of the prodrug and drug is assessed by the concentration required to inhibit cell growth by 50 % (IC<sub>50</sub>).

Treatment of non-transfected or transfected HCT116 cells with the drug results in an IC<sub>50</sub> of approximately 1 µM. In contrast, the PGP prodrug results in an IC<sub>50</sub> of approximately 30 200 µM on non-transfected cells and approximately 1 µM on transfected cells. These results demonstrate that the transfected cells which express the antibody-CPG2 fusion protein can convert the PGP prodrug into the more potent active drug while non-transfected HCT116 cells

are unable to convert the prodrug. Consequently the transfected HCT116 cells are over 100 fold more sensitive to the PGP prodrug in terms of cell killing compared to the non-transfected HCT116 cells. (See Example 1 j) for issues involving possible folic acid depletion in cells).

These studies demonstrate that transfecting tumour cells with a gene for an antibodyenzyme fusion protein can lead to selective tumour cell killing with a prodrug.

#### Example 3

Anti-tumour activity of PGP prodrug in HCT116 tumours expressing the antibody-10 CPG2 fusion protein.

The anti-tumour activity *in vivo* of the PGP prodrug in HCT116 tumours expressing the antibody-CPG2 fusion protein can be demonstrated as follows. HCT116 tumour cells transfected with the antibody-CPG2 fusion protein gene or control non-transfected HCT116 tumour cells are injected subcutaneously into athymic nude mice (10<sup>7</sup> tumour cells per mouse). When the tumours are 5-7 mm in diameter the PGP prodrug is administered i.p. to the mice (3 doses at hourly intervals over 2 h in dose ranges of 5-25 mg kg<sup>-1</sup>). The anti-tumour effects are judged by measuring the length of the tumours in two directions and calculating the tumour volume using the formula:

Volume =  $\Pi/6 \times D^2 \times d$ 

20 where D is the larger diameter and d is the smaller diameter of the tumour.

Tumour volume is expressed relative to the tumour volume at the time the PGP prodrug is administered. The anti-tumour activity is compared to a control group receiving either transfected or non-transfected tumour cells and PBS (170 mM NaCl, 3.4 mM KCl, 12 mM Na<sub>2</sub>HPO<sub>4</sub> and 1.8 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.2) instead of the PGP prodrug.

Administration of PGP to HCT116 tumours established from transfected HCT116 cells results in a significant anti-tumour effect as judged by the PGP treated tumours decreasing in size compared to the PBS treated tumours and it taking a significantly longer time for the PGP treated tumours to reach 4 times their initial tumour volume compared to PBS treated tumours. In contrast, administration of PGP to HCT116 tumours established from non-transfected cells resulted in no significant anti-tumour activity.

Similar studies can be used to demonstrate that the antibody-enzyme gene delivered in an appropriate vector to established HCT116 tumours produced from non-transfected

HCT116 cells when used in combination with the PGP prodrug can result in significant anti-tumour activity. Thus non-transfected HCT116 cells are injected into athymic nude mice (1 X 107 tumour cells per mouse) and once the tumours are 5-7 mm in diameter the vector containing the antibody-enzyme fusion protein gene is injected intra-tumourally. After 1-3 days to allow the antibody-enzyme fusion protein to be expressed by and bind to the HCT116 tumour cells, the PGP prodrug is administered as described above. This results in significant anti-tumour activity compared to control mice receiving PBS instead of PGP prodrug.

## Example 4

# 10 Improved Transfection of Adherent Cell lines Using supplemented FAS media and/or V-79 Feeder Cells

It was foreseen that *in vitro* expression of CPG2 and CPG2 fusion proteins in mammalian cells may degrade media folates leading to slow cell growth or cell death. FAS (folinic acid supplemented) media described herein was developed for CPG2 and CPG2 fusion protein expressing cell lines in order to better support the growth of such cell lines.

In preparation for transfection, adherent cell lines were cultured in normal DMEM media and passaged at least three times before transfection. V-79 (hamster lung fibroblast, obtained from MRC Radiobiology Unit, Harwell, Oxford, United Kingdom) feeder cells were cultured in normal DMEM media and passaged three times before use. For the transfection, a viable count (using a haemocytometer/trypan blue staining) of the adherent cells was made and the cells plated out at 2 x10<sup>5</sup> cells per well into a 6 well plate (Costar 3516) and left for 18-24 hours for the cells to re-adhere.

For each individual transfection, 20µl of LIPOFECTIN<sup>TM</sup> was added to 80µl serum free medium and left at room temperature for 30 minutes. Plasmid DNA (2µg) of interest was added to 100µl serum free medium and subsequently added to the LIPOFECTIN<sup>TM</sup> mix and left for a further 15minutes. The individual 6 well plates were washed with 2 ml serum free medium per well to remove any serum and replaced with 800 µl of fresh serum free medium. The 200µl DNA / LIPOFECTIN<sup>TM</sup>/serum free medium mixes which had been previously prepared were then added to each well of cells. The plates were incubated at 37° for 5 hours, the media removed and 2 ml of fresh normal media added and incubated for a further 48 hours. The transfected cells in the 6 well plate were scraped free, the cell suspension removed and centrifuged. All the supernatant was removed and the cell pellet resuspended in 20 ml of

the appropriate fresh growth media (e.g. FAS DMEM media) containing the appropriate selective agent for the transfected DNA (e.g. G418). Aliquots (200  $\mu$ l) were plated per well into a 96 well plate (1.25  $\times$ 10<sup>4</sup> cells per well).

To enhance clone expansion, fibroblast feeder cells may be added to the transfected 5 cells. Semi-confluent V-79 feeder cells were trypsinised and a viable count performed. The cells were resuspended to 1x106 cells /ml in a sterile glass container, irradiated using a Caesium source by exposure to 5000 rads over 12 minutes. The cells can then be stored at 4° for 24-48 hours (irradiated cells are metabolically active but will not divide, and so can act as "feeders" for other cells without contaminating the culture). The feeder cells should be plated out at 4x104 cells per well in a 96 well plate to produce a confluent layer for the emerging recombinant clones. Feeder cells initially adhere to the plate but with time detach and float off into the media, leaving the any recombinant clone still attached to the well. Media changes (200µl at time) are performed twice weekly to remove floating cells and replenish media. Colonies were allowed to develop for 10-14 days, then the supernatant screened by standard 15 ELISA assay for fusion protein secretion.

To measure the expression rate in the case of the (A5B7-CPG2)<sub>2</sub> fusion gene constructs, recombinant cells were seeded out at 1 x 10<sup>6</sup> in 10 ml fresh normal culture media for exactly 24 hours. The supernatant was then removed, centrifuged to remove cell debris and assayed for fusion protein and enzyme activity by the ELISA and HPLC methods described 20 above. The results for a number of recombinant (A5B7-CPG2)<sub>2</sub> fusion protein cell lines are shown below.

Cell Line	Clone	ng/106cells/24h
HCT 116	F7	6550
	C12	3210
HCT 116	F6	15560
	C1	6151
	В3	4502
	A8	4650
	D5	630
	Н9	610

	G11	2081
	H4	2380
	A4	1634
LoVo	B9	8370
	C1	7350
	F12	2983
	C7	10770
	G10	4140
Colo 320DM	В3	10540
	G4	4720
	B9	885
	B10	3090
	F12	35660

Construction of a stable inducible (A5B7-CPG2)<sub>2</sub> fusion protein expressing tumour cell line

# 5 a) Construction of an inducible fusion protein expression vector

To facilitate expression from a single inducible mammalian cell promoter, an IRES (Internal Ribosome Entry Site; see Y. Sugimoto *et al.*, Biotechnology (1994), **12**, 694-8) based version of the (A5B7-CPG2)<sub>2</sub> fusion protein was constructed. Construct pNG3 pNG3/A5B7VK-HuCK-NEO (A5B7 chimaeric light chain; described in Example 1b above)

- 10 was used as a template for amplification of the light chain gene. The gene was amplified using oligonucleotides CME 3153 and CME 3231 (SEQ ID NOS 19 and 20). A PCR product of the expected size (approximately 700 b.p.) was purified. This product was then digested using the restriction enzymes EcoRI and BamHI and subsequently purified. The fragment was cloned into the Bluescript™ KS+ vector (prepared to receive the fragment by
- digestion with the same restriction enzymes, EcoRI and BamHI) after which the DNA was dephosphorylated and the larger vector band purified. The similarly restricted PCR fragment ligated in to the prepared vector and the ligation mix was transformed into E. coli. DNA was prepared from the clones obtained and analysed by restriction digestion to check for insertion of PCR fragment. Appropriate clones were sequenced to confirm the gene sequence. A

number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation A5B7 Bluescript<sup>TM</sup>.

In a similar manner, the chimaeric A5B7 heavy chain was amplified by PCR from the plasmid pNG4/A5B7VH-IgG2CH1/CPG2 R6 (described in Example 1e above) using 5 oligonucleotides CME 3151 and CME 3152 (SEQ ID NOS 21 and 22). A PCR reaction product of the expected size (approximately 1800 b.p.) was purified. This product was then digested using the restriction enzymes BamHI and Xba I after which the fragment band was purified. The fragment was also cloned into the Bluescript<sup>TM</sup> KS+ vector which had been prepared to receive the above fragment by digestion with the same restriction enzymes,

- 10 BamHl and XbaI, after which the DNA was dephosphorylated and the larger vector band was purified. The similarly restricted PCR fragment was ligated in to the prepared vector and the ligation mix was transformed into E. coli. DNA was prepared from the clones obtained and analysed by restriction digestion to check for insertion of PCR fragment. Appropriate clones were sequenced to confirm the gene sequence. A number of the clones with the correct
- 15 sequence were obtained and one of these clones was given the plasmid designation Bluescript™ Fd-CPG2 R6.

The IRES sequence was sourced from the vector pSXLC (described in Y. Sugimoto et al. Biotechnology (1994), 12, 694-8, and obtained from the authors). The IRES sequence was excised by digestion with the restriction enzymes BamHI and Ncol. A band of the

- 20 expected size (approximately 500 b.p.) was purified and ligated into the Bluescript™ Fd-CPG2 R6 plasmid (which had previously been prepared by restriction with the same enzymes). The ligation mix was transformed into E. coli and DNA was prepared from the clones obtained. The DNA was analysed by restriction digestion to check for insertion of the fragment and appropriate clones were subsequently sequenced to confirm the gene sequence.
- 25 A number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation **Bluescript<sup>TM</sup> IRES Fd-CPG2 R6**.

To facilitate later cloning steps, it was necessary to delete the Xba I site which had been carried over in the IRES fragment. This was performed by PCR mutagenesis with the oligonucleotide primers CME 3322 and CME 3306 (SEQ ID NOS: 23 and 24) and the

30 Bluescript<sup>TM</sup> IRES Fd-CPG2 R6 as template DNA. A PCR reaction product of the expected size (approximately 500 b.p.) was purified, digested with the restriction enzymes BamHI and NcoI and ligated into the Bluescript<sup>TM</sup> IRES Fd-CPG2 R6 plasmid (which had previously

been prepared by restriction with the same restriction enzymes). The ligation mix was transformed into *E. coli* and DNA was prepared from the clones obtained. The DNA was analysed by restriction digestion to check for insertion of the fragment and appropriate clones were subsequently sequenced to confirm the gene sequence. A number of the clones with the 5 correct sequence were obtained and one of these clones was given the plasmid designation Bluescript<sup>TM</sup> IRES Fd-CPG2 R6-Xba del.

The A5B7 chimaeric light chain fragmentwas excised from the A5B7 Bluescript<sup>TM</sup> plasmid by digestion with the restriction enzymes EcoR1 and BamHI. A band of the expected size (approximately 700 b.p.) was purified, ligated into the appropriately prepared Bluescript 10 IRES Fd-CPG2 R6-Xba del plasmid and the ligation mix was transformed into E. coli. DNA was prepared from the clones obtained and analysed by restriction digestion to check for insertion of the fragment. Appropriate clones were subsequently sequenced to confirm the gene sequence. A number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation Bluescript<sup>TM</sup> A5B7 IRES Fd-CPG2 R6-Xba del. The complete IRES based A5B7 chimaeric fusion protein gene sequence is shown in SEQ ID NO: 52.

The IRES based A5B7 chimaeric fusion protein gene was then transferred to a tetracycline regulated expression vector. Vectors for the Tet On gene expression system were obtained from Clontech. The Tetracycline switchable expression vector pTRE (otherwise known as pHUD10-3, see Gossen et al. (1992), PNAS, 89, 5547-51) was prepared to accept the IRES based fusion protein cassette by digestion with the restriction enzymes EcoR1 and XbaI, dephosphorylated and the larger vector band purified. The IRES gene cassette was excised from the Bluescript<sup>TM</sup> A5B7 IRES Fd-CPG2 R6-Xba del plasmid using the same restriction enzymes. The approximately 3000 b.p. fragment obtained was ligated in to the prepared vector and the ligation mix was transformed into E. coli. DNA was prepared from the clones obtained and analysed by restriction digestion to check for insertion of PCR fragment. Appropriate clones were subsequently sequenced to confirm the gene sequence. A number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation pHUD10-3/A5B7 IRES Fd-CPG2 R6.

# 30 b) Construction of a stable inducible fusion protein expressing cell line The standard lipofection transfection methodology (as described previously but

without the use of feeder cells) was used to produce recombinant HCT116 tumour cell lines.

A co-transfection using 1 $\mu$ g of the pHUD10-3/A5B7 IRES Fd-CPG2 R6 plasmid and 1 $\mu$ g of the pTet-On transactivator expressing plasmid (from the Clontech kit) was performed and positive clones selected using FAS media containing 750  $\mu$ g G418/ml .

## c) Induction studies of recombinant HCT116 inducible cell lines

The clone cultures obtained were split in to duplicate 48 well plates, each containing lx 106 cells. The cells were grown for 48 h with one of the plates induced with 2 μg/ml doxycycline and the other acting as an non-induced control. Expression of the (A5B7-CPG2)<sub>2</sub> fusion protein in the cell supernatant was tested using the ELISA/ Western blot assays described in Example 1g. The results indicated that induction of fusion protein from the inducible cell line by use of doxycycline could be clearly demonstrated, for example one of the clones obtained (F11), the induced cells produced 120 ng/ml of fusion protein in the supernatant whereas the non-induced cells produced only background levels of fusion protein (below 1 ng/ml).

## 15 Example 6

## Cell based ELISA assay of secreted fusion protein material

Cells were seeded into 96 well plates (Becton Dickinson Biocoat<sup>TM</sup> poly-<u>D</u>-Lysine, 35-6461) at a density of  $1x10^4$  cells per well in  $100 \mu l$  normal culture media and left about 40 h at 37°.  $100 \mu l$  of 6 % formaldehyde was diluted in DMEM and left for 1 hour at 4°. Plates 20 were centrifuged and washed 3 times in PBS containing 0.05 % Tween<sup>TM</sup> by immersion soaking (first two washes for 2 minutes and the final wash for 5 minutes).

100μl of doubling dilutions of cell culture supernatant containing fusion protein or chimeric A5B7 anti-CEA were added to each well as appropriate and the plates incubated overnight at 4°. The plates were washed as described above and, in the case of chimaeric fusion proteins, 100μl of 1:1000 dilution of HRP labelled anti-human kappa antibody (Sigma A-7164) was added and incubated for 2 hours at room temperature (an anti-CPG2 detection methodology can be used in the case of murine scFv fusion proteins). The plates were washed as described above and HRP detected using OPD substrate (Sigma P-8412). Colour was allowed to develop for about 5 min, stopped with 75 μl per well of 2M H<sub>2</sub>SO<sub>4</sub> and OD read at 490 nm.

In the case of the (A5B7-CPG2)<sub>2</sub> fusion protein, material was produced in the

supernatant from recombinant Colo320DM tumour cells (CEA-ve). The fusion protein content was measured by use of the CEA ELISAs described above. Increasing amounts of fusion protein were added to a number of CEA negative cell lines and the CEA positive LoVoparental line. The results shown in Figure 3 clearly show that only the CEA positive line 5 shows increased levels of binding with increasing amounts of added fusion protein whereas the CEA negative cell lines show only constant background binding levels throughout. This clearly demonstrates that the fusion protein specifically binds and is retained on CEA positive

## 10 Example 7

Lovo cells.

# Recombinant LoVo tumour cells expressing antibody-enzyme fusion protein exhibit retention of the fusion protein on the cell surface

LoVo colorectal tumour cells transfected with the (A5B7-CPG2)<sub>2</sub> fusion protein gene have been shown both to secrete and to retain the fusion protein on their cell surface.

- 15 This can be demonstrated by comparing parental and recombinant fusion protein expressing LoVo cells under the conditions set out in the cell based ELISA assay of secreted fusion protein (Figure 4). On development of the colour reaction it could be seen that the recombinant LoVo cells had retained the expressed fusion protein (by showing a high level of colour). In control experiments, using Colo320DM fusion protein expressing cells, the assay
- 20 showed some retention of the expressed fusion protein (probably non-specific) and the parental LoVo cells only exhibited background activity. Positive controls in which CEA binding antibody was added to test recombinant fusion protein expressing tumour cells and to the parental LoVo controls resulted in a signal being obtained from the parental LoVo (thus demonstrating that CEA was present on the parental cells) but no increased signal from the
- 25 Colo320DM (CEA negative). The recombinant LoVo cells still gave such a strong initial signal that the added antibody made little difference to the overall signal obtained, which was considerably higher than any of the control experiments. Thus it appears that anti-CEA antibody enzyme-CPG2 fusion protein secreted from CEA positive tumour cell lines bind to the surface of the cells (via CEA) whereas the same protein expressed from CEA negative
- 30 tumours shows no such binding.

LoVo tumour cells expressing the antibody-enzyme fusion protein are selectively killed in vitro by a prodrug.

LoVo colorectal tumour cells, transfected with the (A5B7-CPG2)<sub>2</sub> fusion protein gene, 5 can be selectively killed by a prodrug that is converted by CPG2 enzyme into an active drug.

To demonstrate this control non-transfected LoVo cells or LoVo cells transfected with an antibody-CPG2 fusion protein gene are incubated with either the prodrug, 4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl-L-glutamic acid (PGP; Blakey et al, (1995) Br. J. cancer 72, p1083) or the corresponding drug released by CPG2, 4-[N,N-bis(2-chloroethyl)amino] phenol as described in Example 2 with HCT116 cells.

The transfected cells which express the antibody-CPG2 fusion protein can convert the PGP prodrug into the more potent active drug while non-transfected LoVo cells are unable to convert the prodrug.

These studies demonstrate that transfecting tumour cells with a gene for an antibody-15 enzyme fusion protein can lead to selective tumour cell killing with a prodrug.

## Example 9

# Establishment of fusion protein expressing LoVo tumour xenografts in athymic mice

Recombinant LoVo fusion protein (A5B7-CPG2)<sub>2</sub> expressing tumour cells or mixes of recombinant and parental LoVo cells were injected subcutaneously into athymic nude mice (10<sup>7</sup> tumour cells per mouse). The tumour growth rates for both 100 % recombinant and 20 %: 80 % mixes of recombinant:parental LoVo cells were compared to those of parental cell only tumours. No significant differences were seen in the observed growth curves obtained showing no corrections were required during comparisons between the cell lines. The tumour growth rates observed showed that in each case for the xenograft tumours to reach a size of 10 x 10 mm takes about 12 days.

# Determination of enzyme activity in tumour xenograft samples

To act as a standard for the assay, a CPG2 enzyme standard curve was prepared in 20 % homogenate of normal tumour (parental cell tumour). Subsequent dilutions of samples 5 were made in the same 20 % homogenate of normal tumour.

Excised tumour tissue is removed from -80° storage (previously flash frozen in liquid nitrogen) and allowed to thaw. Any residual skin tissue was removed before the tumour was cut up in to small fragments with a scalpel. The tumour tissue was transferred to a preweighed tube and the weight of tumour tissue measured. PBS containing 0.2 mM ZnCl<sub>2</sub> solution was added to each tumour sample to give a 20 % (w/v) mix, homogenised and placed on ice. Dilutions of sample tumours (in 20% normal tumour homogenate) were prepared e.g. neat, 1/10, 1/20 and 1/40.

For the standard curve, dilutions of CPG2 enzyme were made to the following concentrations to a final volume of 400 µl. Similarly, 400 µl of each of the recombinant 115 tumour sample dilutions were also prepared. After equilibration to 30°, 4 µl of 10 mM methotrexate (MTX) solution was added. The reaction was stopped after exactly 10 minutes by adding 600 µl ice cold methanol + 0.2 % TFA, centrifuged and the supernatant collected. The substrate and product in the supernatant were then separated by HPLC (using a Cation Exchange Column, HICROM™ S5SCX-100A, mobile phase = 60 % methanol, 40 % 60 mM 20 ammonium formate/ 0.1 % TFA, detection 300 nm). To calculate enzyme activity in the tumour tissue, the standard curve was plotted as units of area of methotrexate metabolite (the standards are such that only 20-30 % of the substrate is metabolised so ensuring this is not rate limiting). The test samples were analysed by comparing the unit area of metabolite against the standard curve and then multiplying by the dilution factor. Finally, making the 25 working assumption that 1 ml= 1 g the results were multiplied by 5 (as the samples were originally diluted to a 20% homogenate).

Results obtained with 20 % recombinant: 80 % parental LoVo cells expressing (A5B7 Fab-CPG2)<sub>2</sub> fusion protein showed the following results: tumours taken at day 5 had an average enzyme activity = 0.26 U/g (range between 0.18-0.36 U/g) and at day 12 had an 30 average enzyme activity = 0.65 U/g (range between 0.19-1.1 U/g).

## Determination enzyme activity in plasma samples

To act as a standard for the assay, a CPG2 enzyme standard curve was prepared in 20 % normal plasma to the following concentrations: 0.2, 0.4, 0.6, 0.8 and 1.0 U/ml. Similarly all test plasma samples were also diluted to 20 % normal plasma. Further dilutions of these samples e.g. neat 1/10, 1/20 and 1/50 were also made using 20 % normal serum. 200 μl aliquots of each CPG2 standard and test sample dilutions were equilibrated to 30°. 2 μl of 10 mM MTX was added to each of the tubes and mixed well. to 30°. The reaction was stopped after exactly 10 minutes (to increase the sensitivity of the assay the incubation time can be 10 increased to 30 minutes) by adding 500 μl ice cold methanol + 0.2 % TFA and assayed for product using HPLC detection as described above in Example 10.

No activity was seen in the plasma except in the rare cases when the level of enzyme activity in the tumour exceeded 2.0 U/g, in which case the plasma enzyme levels were measured in the range of 0.013 to 0.045 U/ml.

15

### Example 12

# Anti-tumour activity of PGP prodrug in LoVo tumours expressing the antibody-CPG2 fusion protein.

Recombinant LoVo (A5B7-CPG2)<sub>2</sub> fusion protein expressing tumour cells or mixes of recombinant and parental LoVo cells were injected subcutaneously into athymic nude mice as described in Example 9.

When the tumours are 5-7 mm in diameter the PGP prodrug is administered i.p. to the mice (3 doses in DMSO/ 0.15 M sodium bicarbonate buffer at hourly intervals over 2 h in dose ranges of 40-80 mg kg<sup>-1</sup>).

Anti-tumour effects are judged by measuring the length of the tumours in two directions and calculating the tumour volume using the formula

Volume = 
$$\Pi/6 \times D^2 \times d$$

where D is the larger diameter and d is the smaller diameter of the tumour. Tumour volume may be expressed relative to the tumour volume at the time the PGP prodrug is administered or alternatively the median tumour volumes may be calculated. The anti-tumour activity is compared to control groups receiving either transfected or non-transfected tumour cells and

buffer without PGP prodrug.

Administration of PGP to LoVo tumours established from recombinant LoVo cells or recombinant Lovo/Parental LoVo cell mixes results in a significant anti-tumour effect as shown by the PGP treated tumours decreasing in size compared with controls and it taking a 5 significantly longer time for the PGP treated tumours to reach 4 times their initial tumour volume compared with controls (Figure 5). Administration of PGP to LoVo tumours established from non-transfected cells resulted in no significant anti-tumour activity.

Similar studies can be used to demonstrate that the antibody-enzyme gene delivered in an appropriate gene delivery vector to established LoVo tumours produced from non-10 transfected parental LoVo cells when used in combination with the PGP prodrug can result in significant anti-tumour activity. Thus non-transfected LoVo cells are injected into athymic nude mice (1 X 107 tumour cells per mouse) and once the tumours are 5-7 mm in diameter the vector containing the antibody-enzyme fusion protein gene is injected intra-tumourally. After 1-3 days to allow the antibody-enzyme fusion protein to be expressed by, and bind to, the LoVo tumour cells, the PGP prodrug is administered as described above. This results in significant anti-tumour activity compared with controls.

#### Example 13

## Construction of an (806.077 Fab-CPG2)2 fusion protein

The construction of a (806.077 Fab-CPG2)<sub>2</sub> enzyme fusion was planned with the aim of obtaining a bivalent human carcinoembryonic antigen (CEA) binding molecule which also exhibits CPG2 enzyme activity. To this end the initial construct was designed to contain an 806.077 antibody heavy chain Fd fragment linked at its C-terminus via a flexible (G4S)<sub>3</sub> peptide linker to the N-terminus of the CPG2 polypeptide (as shown in Figure 1 but 25 substituting 806.077 in place of A5B7).

The antibody 806.077 (described in International Patent Application WO 97/42329, Zeneca Limited) binds with a very high degree of specificity to human CEA. Thus the 806.077 antibody is particularly suitable for targeting colorectal carcinoma or other CEA antigen bearing cells.

In general, antibody (or antibody fragment)-enzyme conjugate or fusion proteins should be at least divalent, that is to say capable of binding at least 2 tumour associated antigens (which may be the same or different). In the case of the (806.077 Fab-CPG2), fusion

protein, dimerisation of the enzyme component takes place (after expression, as with the native enzyme) thus forming an enzymatic molecule which contains two Fab antibody fragments (and is thus bivalent with respect to antibody binding sites) and two molecules of CPG2 (Figure 2a).

## 5 a) Cloning of the 806.077 antibody genes

Methods for the cloning and characterisation of recombinant murine 806.077 F(ab')<sub>2</sub> antibody have been published (International Patent Application WO 97/42329, Example 7). Reference Example 7.5, describes cloning of the 806.077 antibody variable region genes into Bluescript<sup>TM</sup> KS+ vectors. These vectors were subsequently used as the source of the 806.077 variable region genes for the construction of 806.077 chimaeric light and heavy chain Fd genes.

## b) Chimaeric 806.077 antibody vector constructs

International Patent Application WO 97/42329, Example 8 describes the cloning of the 806.077 chimaeric light and heavy chain Fd genes in the vectors pNG3-Vkss-HuCk-NEO (NCIMB deposit no. 40799) and pNG4-VHss-HuIgG2CH1' (NCIMB deposit no. 40797) respectively. The resulting vectors were designated pNG4/VHss806.077VH-IgG2CH1' (806.077 chimaeric heavy chain Fd') and pNG3/VKss806.077VK-HuCK-NEO (806.077 chimaeric light chain). These vectors were the source of the 806.077 antibody genes for the construction of the 806.077 Fab-CPG2 fusion protein.

# 20 c) Construction of the 806.077 heavy chain Fd-CPG2 fusion protein gene

The cloning and construction of the CPG2 gene used are described in Example 1, sections c and d. Similarly, the construction of the pNG4/A5B7VH-IgG2CH1/CPG2 R6 vector, which was used for the construction of the 806.077 heavy chain Fd-CPG2, is described in Example 1, section e. The 806.077 variable heavy chain gene was removed from the

- pNG4/VHss806.077VH-IgG2CH1' vector by digestion with restriction enzymes HindIII and NheI and a band of the expected size (approximately 300 b.p) which contained the variable region gene was purified. The same restriction enzymes (HindII/NheI) were used to digest the vector pNG4/A5B7VH-IgG2CH1/CPG2 R6 in preparation for the substitution of the 806.077 variable region for that of the A5B7 antibody. After digestion, the DNA was
- 30 dephosphorylated then the larger vector band was separated and purified. The similarly restricted variable region gene fragment was then ligated in to this prepared vector and the ligation mix transformed into E. coli. DNA was prepared from the clones obtained and

analysed by restriction digest analysis and subsequently sequenced to confirm the fusion gene sequence. A number of the clones were found to be correct and one of these clones, pNG4/VHss806VH-lgG2CH1/CPG2 R6, was chosen for further work. The sequence of the 806.077 heavy chain Fd-CPG2 fusion protein gene created is shown SEQ ID NOS 25 and 26.

# 5 d) Co-transfection, transient expression and analysis of fusion protein

The plasmids pNG4/VHss806.077VH-IgG2CH1/CPG2 R6 (encoding the antibody chimaeric Fd-CPG2 fusion protein) and pNG3/VHss806.077VK-HuCK-NEO (encoding the antibody chimaeric light chain) were co-transfected into COS-7 cells using a LIPOFECTINTM based procedure described in Example 1f above. Analysis of the fusion protein was

- performed as described in Example 1g. The HPLC based enzyme activity assay clearly showed CPG2 enzyme activity to be present in the cell supernatant and both the anti-CEA ELISA assays exhibited binding of protein at levels commensurate with a bivalent 806.077 antibody molecule. The fact that the anti-CEA ELISA detected with an anti-CPG2 reporter antibody also exhibited clear CEA binding indicated that not only antibody but also antibody-
- 15 CPG2 fusion protein was binding CEA. Western blot analysis with both reporter antibody assays clearly displayed a (806.077 Fab-CPG2)<sub>2</sub> fusion protein subunit of the expected approximately 90 kDa size with only a small amount of degradation or smaller products (such as Fab or enzyme) observable. Since CPG2 is only known to exhibit enzyme activity when it is in a dimeric state it and since only antibody enzyme fusion protein is present, this indicates
- 20 that the 90 kDa fusion protein (seen under SDS/PAGE conditions) dimerises via the natural CPG2 dimerisation mechanism to form a 180 kDa dimeric antibody-enzyme fusion protein molecule (Figure 2a) in "native" buffer conditions. Furthermore, this molecule exhibits both CPG2 enzymatic activity and CEA antigen binding properties which do not appear to be significantly different in the fusion protein compared with enzyme or antibody alone.

# 25 e) Construction of a (806.077 Fab-CPG2)<sub>2</sub> fusion protein coexpression vector for use in transient and stable cell line expression

For a simpler transfection methodology and the direct coupling of both expression cassettes to a single selection marker, a co-expression vector for fusion protein expression was constructed using the existing vectors pNG4/VHss806.077VH-IgG2CH1/CPG2 (encoding the antibody Fd-CPG2 fusion protein) and pNG3/VKss806.077VK-HuCK-NEO (encoding the antibody light chain). The pNG4/VHss806.077VH-IgG2CH1/CPG2 plasmid was first digested with the restriction enzyme Scal, the linear vector band purified, digested with the

restriction enzymes BgIII and BamHI and a desired band (approximately 2700 b.p.) purified. The plasmid pNG3/VKss806.077VK-HuCK-NEO was digested with the restriction enzyme BamHI after which the DNA was dephosphorylated and the vector band purified. The heavy chain expression cassette fragment was ligated in to the prepared vector and the ligation mix transformed into *E. coli*. The orientation was checked by a variety of restriction digests and clones selected which had the heavy chain cassette in the same direction as that of the light chain. This plasmid was termed pNG3-806.077-CPG2/R6-coexp.-NEO.

## Example 14

# 10 Construction of a (55.1 scFv-CPG2)<sub>2</sub> fusion protein

The 55.1 antibody, described in the United States Patent 5,665,357, recognises the CA55.1 tumour associated antigen which is expressed on the majority of colorectal tumours and is only weakly expressed or absent in normal colonic tissue. The determination of the 55.1 heavy and light chain cDNA sequences is described in Example 3 of the aforementioned 15 US patent. A plasmid expression vector allowing the secretion of antibody fragments into the periplasm of *E.coli* utilizing a single pelB leader sequence (pICI266) has been deposited as accession number NCIMB 40589 on 11Oct93 under the Budapest Treaty at the National Collections of Industrial and Marine Bacteria Limited (NCIMB), 23 St. Machar Drive, Aberdeen, AB2 1RY, Scotland, U.K. This vector was modified as described in Example 3.3a of United States Patent 5,665,357 to create pICI1646; this plasmid was used for cloning of various 55.1 antibody fragments as described in further subsections of Example 3, including the production of a 55.1 scFv construct which was designated pICI1657.

The pICI1657 (otherwise known as pICI-55.1 scFv) was used as the starting point for the construction of the (55.1 scFv-CPG2)<sub>2</sub> fusion protein. The 55.1 scFv gene was amplified using the oligonucleotides CME 3270 and CME 3272 (SEQ ID NOS: 27 and 28 respectively) and the plasmid pICI1657 as the template DNA. The resulting PCR product band of about 790 b.p. was purified. Similarly the pNG4/A5B7VH-IgG2CH1/CPG2 R6 plasmid described in Example 1e above was used as the template DNA in a standard PCR reaction to amplify the CPG2 gene using the oligonucleotide primers CME 3274 and CME 3275 (SEQ ID NOS: 29 and 30 respectively). The expected PCR product band of about 1200 b.p. was purified.

A further PCR reaction was performed to join (or splice) the two purified PCR reaction products together. Standard PCR reaction conditions were used using varying

amounts (between 0.5 to 2 μl) of each PCR product but utilising 25 cycles (instead of the usual 15 cycles) with the oligonucleotides CME 3270 and CME 3275 (SEQ ID NOS: 27 & 30). A reaction product of the expected size (approximately 2000 b.p.) was excised, purified and eluted in 20 μl H<sub>2</sub>O, digested using the restriction enzyme EcoRI and purified. The 5 vector pNG4/VHss806.077VH-IgG2CH1/CPG2 was prepared to receive the above PCR product by digestion with restriction enzyme EcoRI, dephosphorylated, the larger vector band separated from the smaller fragment and purified. The similarly restricted PCR product was ligated in to the prepared vector and the ligation mix transformed into *E. coli*. DNA was prepared from the clones obtained and analysed by HindIII/NotI restriction digestion to check for correct fragment orientation and appropriate clones subsequently sequenced to confirm the fusion gene sequence. A number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation pNG4/55.1scFv/CPG2 R6. The DNA

## 15 Example 15

# Modification of the plasmid pNG4/55.1scFv/CPG2 R6 to facilitate scFv gene exchange

and amino acid sequences of the fusion protein are shown in SEQ ID NOS: 31 and 32.

During the construction of pNG4/55.1scFv/CPG2 R6 a unique BspEI (isoschizomer of AccIII) was introduced into the flexible (G<sub>4</sub>S)<sub>3</sub> linker coding sequence, situated between the antibody and CPG2 genes. To facilitate cloning of alternative scFv constructs the EcoRI 20 site 3' of the CPG2 gene in the pNG4/55.1scFv/CPG2 R6 was deleted in order to enable insertion of alternative scFv antibody genes in frame, both behind the plasmid signal sequence and 5' of the CPG2 gene, via a EcoRI/BspEI fragment cloning. This modification was achieved by PCR mutagenesis in which first the pNG4/55.1scFv/CPG2 R6 was amplified using oligonucleotides CME 3903 and CME 3906 (SEQ ID NOS: 33 and 34 respectively).

25 Secondly, the pNG4/55.1scFv/CPG2 R6 was again amplified but using oligonucleotides CME 4040 and CME 3905 (SEQ ID NOS: 35 and 36 respectively). The first expected PCR product band of about 420 b.p. was purified. The second PCR reaction was similarly treated and the expected PCR product band of about 450 b.p. purified.

A further PCR reaction was performed to join (or splice) the two purified PCR 30 reaction products together. Standard PCR reaction conditions were used using varying amounts (between 0.5 to 2 µl) of each PCR product but utilising between 15 and 25 cycles

with oligonucleotides CME 3905 and CME 3906 (SEQ ID NOS: 36 & 34). A reaction product of the expected size (approximately 840 b.p.) was purified, digested using the restriction enzymes NotI and XbaI and the expected fragment band of ca.460 b.p. was purified.

The original pNG4/55.1scFv/CPG2 R6 was prepared to receive the above PCR product by digestion with restriction enzymes NotI and XbaI, dephosphorylated and the larger vector band separated from the smaller fragment. The vector band was purified and subsequently the similarly restricted PCR product was ligated in to the prepared vector and the ligation mix transformed into *E. coli*. DNA was prepared from the clones obtained and analysed by EcoRI restriction digestion to check for insertion of the modified fragment and appropriate clones subsequently sequenced to confirm the sequence change. A number of clones with the correct sequence were obtained and one of these clones was given the plasmid designation pNG4/55.1scFv/CPG2 R6/del EcoRI. This mutation removes the EcoRI site which was 3' of the CPG2 gene and simultaneously introduces an additional stop codon. The DNA sequence of the fusion protein gene up to, and including the two stop codons, are shown in SEQ ID NO: 37.

#### Example 16

## Construction of an 806.077 scFv antibody gene

The 806.077 scFv was created using vectors pNG4/VHss806.077VH-IgG2CH1' and pNG3/VKss806.077VK-HuCK-NEO which are sources for 806.077 VH and VK variable region genes. The 806.077 VH gene was amplified from the pNG4/VHss806.077VH-IgG2CH1' plasmid using standard PCR conditions with the oligonucleotides CME 3260 and CME 3266 (SEQ ID NOS: 39 and 40 respectively). The 806.077 VK was amplified from the pNG3/VKss806.077VK-HuCK-NEO plasmid using oligonucleotides CME 3262 and CME 3267 (SEQ ID NOS: 41 and 42 respectively). The VH and VK PCR reaction products were purified.

A further PCR reaction was performed to join (or splice) the two purified PCR reaction products together. Standard PCR reaction conditions were used using varying amounts (between 0.5 to 2 µl) of each PCR product but utilising between 15 and 25 cycles with the flanking oligonucleotides oligonucleotides CME 3260 and CME 3262 (SEQ ID NOS: 39 & 41). A reaction product of the expected size (approximately 730 b.p.) was

purified, digested using the restriction enzymes NcoI and XhoI and an expected fragment band of about 720 b.p. purified.

The pICI1657 plasmid (otherwise known as pICI-55.1 scFv) had been further modified by the insertion of a double stranded DNA cassette produced from the two 5 oligonucleotides CME 3143 and CME 3145 (SEQ ID NOS: 45 and 46) between the existing XhoI and EcoR restriction sites by standard cloning techniques to create the vector pICI266-55.1 scFv tag/his (the DNA sequence of the resulting 55.1 scFv tag/his gene is shown in SEQ ID NO: 47). This vector was prepared to receive the above PCR product by digestion with restriction enzymes NcoI and XhoI, dephosphorylated and the larger vector band separated 10 from the smaller fragment. The vector band was purified and subsequently the similarly restricted PCR product was ligated in to the prepared vector and the ligation mix transformed into E. coli. DNA was prepared from the clones obtained and analysed by EcoRI restriction digestion to check for insertion of the modified fragment and appropriate clones subsequently sequenced to confirm the sequence change. A number of the clones with the correct sequence 15 were obtained and one of these clones was given the plasmid designation pICI266/806IscFvtag/his (alternatively known as pICI266-806VH/VLscFvtag/his). The DNA and protein sequences of the 806I scFvtag/his gene are shown in (SEQ ID NOS: 25 and 26).

## 20 Example 17

## Construction of an (806.077 scFv-CPG2)2 fusion protein

The pICI266/806IscFvtag/his plasmid was used as the source for the 806scFv. The gene was amplified using oligonucleotides CME 3907 and CME 3908 (SEQ ID NOS: 48 and 49) and a band of the expected size purified. This fragment was then digested using the restriction enzymes EcoRI and BspEI after which an expected fragment band of about 760 b.p. was purified.

The pNG4/55.1scFv/CPG2 R6/del EcoRI plasmid was prepared to receive the above fragment by digestion with restriction enzymes EcoRI and BspEI, dephosphorylated and the larger vector band separated from the smaller fragment. The vector band was purified and subsequently the similarly restricted fragment ligated in to the prepared vector and the ligation mix was transformed into E. coli.. DNA was prepared from the clones obtained and analysed by EcoRI restriction digestion to check for insertion of the modified fragment. Appropriate

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clones were subsequently sequenced to confirm the gene sequence. A number of the clones with the correct sequence were obtained and one of these clones was given the plasmid designation pNG4/806IscFv/CPG2 R6/del EcoRI. The DNA and protein sequence of the fusion protein gene 806IscFv/CPG2 R6 are shown in (SEQ ID NOS: 50 and 51).

5

## Example 18

# Co-transfection, transient expression of antibody-CPG2 fusion proteins

As described in Example 1f, plasmids encoding other fusion protein variants can be transfected using the given standard conditions in order to obtain transient expression of their encoded fusion protein from COS7 cells. In the case of (Fab-CPG2)<sub>2</sub> fusion proteins both cotransfection of appropriate plasmids or transfection of co-expression proteins can be performed. Similarly, the single expression plasmids of (scFv-CPG2)<sub>2</sub> fusion proteins can be also be transfected by the same protocol. In each case a maximum total of 4 mg DNA are used in an individual transfection.

15

#### Example 19

## Gene switches for protein expression

As described in Example 1 j, the use of tightly controlled but inducible gene switch systems such as the "TET on" or "TET off" (Grossen, M. et al (1995) Science 268: 1766-20 1769) or the ecdysone/ muristerone A (No, D. et al (1996) PNAS 93:3346-3351) may be used for the expression of fusion proteins. Appropriate methodology and cloning strategies as described in Example 5 may be used for antibody Fab-enzyme fusions requiring an IRES sequence for expression. Insertion of the appropriate gene cassette in to the switchable expression vectors may be used if the fusion protein product is a single polypeptide chain 25 such as in scFv-enzyme constructs.

#### Example 20

# Determination of the properties of COS7 cell secreted antibody-enzyme fusion proteins

The COS7 cell supernatant material can be analysed for the presence of antibody

30 fusion proteins as described in Example 1g. Similarly the use of expressed fusion protein and

CPG2 prodrug in an *in vitro* cytotoxicity assay can be performed as previously described in

Example 1h. The HPLC based enzyme activity assay can show CPG2 enzyme activity to be

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present in the cell supernatant and anti-CEA ELISA can be detected with an anti-CPG2 reporter antibody to confirm binding of protein at levels commensurate with a bivalent A5B7 antibody molecule and also to demonstrate that antibody-CPG2 fusion protein (not only just the antibody component) is binding CEA.

Western blot analysis with both reporter antibody assays clearly display a fusion protein subunit of the expected size. Since CPG2 is only known to exhibit enzyme activity when it is in a dimeric state it and since only antibody enzyme fusion protein is present, this indicates that the fusion protein (seen under SDS/PAGE conditions) dimerises via the natural CPG2 dimerisation mechanism to form a dimeric antibody-enzyme fusion protein molecule in 10 "native" buffer conditions. Furthermore, this molecule exhibits both CPG2 enzymatic activity and CEA antigen binding properties which do not appear to be significantly different in the fusion protein compared with enzyme or antibody alone. Results obtained from the cytotoxicity assay can demonstrate that antibody-enzyme fusion protein (together with prodrug) causes at least equivalent cell kill and resulted in lower numbers of cells at the end of 15 the assay period than the equivalent levels of A5B7 F(ab')2-CPG2 conjugate (with the same prodrug). Since cell killing (above basal control levels) can only occur if the prodrug is converted to active drug by the CPG2 enzyme (and since the cells are washed to remove unbound protein, only cell bound enzyme will remain at the stage where the prodrug is added). Thus this experiment can demonstrate that at least as much of the (A5B7-CPG2 R6)2 20 fusion protein remains bound compared with conventional A5B7 F(ab)2-CPG2 conjugate as a greater degree of cell killing (presumably due to higher prodrug to drug conversion) occurs.

#### Example 21

In vitro and in vivo determination of the properies of antibody-enzyme fusion proteins expressed from recombinant tumour cells

The construction of fusion protein expressing tumour cell lines can be performed as described in Example 4.

Retention of the fusion protein on the cell surface of recombinant LoVo tumour cells expressing antibody-enzyme fusion protein can be shown using the techniques described in 30 Example 7. Selective killing of cultured LoVo tumour cells transfected with an antibody-CPG2 fusion protein gene by a prodrug that is converted by the enzyme into an active drug

can be demonstrated as described in Example 8. Establishment of antibody-enzyme fusion protein expressing LoVo tumours xenografts in athymic mice can be performed as described in Example 9. Determination of enzyme activity in tumour xenograft samples can also be determined as described in Example 10.

Determination enzyme activity in plasma samples performed as described in Example 11. The anti-tumour activity of PGP prodrug in LoVo tumours expressing the antibody-CPG2 fusion protein can be evaluated using the method described in Example 12.

The results from these experiments can be used to show that the antibody-CPG2 fusion protein secreted from CEA positive tumour cell lines bind to the surface of the cells (via CEA) whereas the same protein expressed from CEA negative tumours shows no such binding.

These results can demonstrate that the transfected cells which express the antibody-CPG2 fusion protein can convert the PGP prodrug into the more potent active drug while non-transfected LoVo cells are unable to convert the prodrug. Consequently the transfected LoVo cells will be over 100 fold more sensitive to the PGP prodrug in terms of cell killing compared to the non-transfected LoVo cells, thus demonstrating that transfecting tumour cells with a gene for an antibody-enzyme fusion protein can lead to selective tumour cell killing with a prodrug.

Administration of PGP to LoVo tumours established from recombinant LoVo cells or recombinant Lovo/Parental LoVo cell mixes can result in a significant anti-tumour effect as judged by the PGP treated tumours decreasing in size compared to the formulation buffer only treated tumours and it taking a significantly longer time for the PGP treated tumours to reach 4 times their initial tumour volume compared with formulation buffer treated tumours. In contrast, administration of PGP to LoVo tumours established from non-transfected cells would result in no significant anti-tumour activity.

Similar studies can be used to demonstrate that the antibody-enzyme gene delivered in an appropriate gene delivery vector to established LoVo tumours produced from non-transfected parental LoVo cells when used in combination with the PGP prodrug can result in significant anti-tumour activity. Thus non-transfected LoVo cells are injected into athymic nude mice (1 X 107 tumour cells per mouse) and once the tumours are 5-7 mm in diameter the vector containing the antibody-enzyme fusion protein gene is injected intra-tumourally. After 1-7 days to allow the antibody-enzyme fusion protein to be expressed by, and bind to, the

LoVo tumour cells, the PGP prodrug is administered as previously described. This results in significant anti-tumour activity compared with control mice receiving formulation buffer instead of PGP prodrug.

#### 5 Example 22

# Preparation of (murine A5B7 Fab-CPG2)2 fusion protein

(Murine A5B7 Fab-CPG2)<sub>2</sub> is expressed from COS-7 and CHO cells essentially as described in part (d) of Example 48 of International Patent Application WO 97/42329 (Zeneca Limited, published 13 November, 1997) by cloning the genes for A5B7 light chain and A5B7 10 Fd linked at its C-terminus via a flexible (G<sub>4</sub>S)<sub>3</sub> peptide linker to CPG2 in the pEE14 co-expression vector.

The murine A5B7 light chain is isolated from pAF8 (described in part g of Reference Example 5 in International Patent Application WO 96/20011, Zeneca Limited). Plasmid pAF8 is cut with EcoRI and the resulting 732 bp fragment isolated by electrophoresis on a 1% agarose gel. This fragment is cloned into pEE14 (described by Bebbington in METHODS: A Companion to Methods in Enzymology (1991) 2, 136-145) similarly cut with EcoRI and the resulting plasmid used to transform *E. coli* strain DH5α. The transformed cells are plated onto L agar plus ampicillin (100 μg/ml). A clone containing a plasmid with the correct sequence and orientation is confirmed by DNA sequence analysis (SEQ ID NO: 57) and the plasmid named pEE14/A5B7muVkmuCK. The amino acid sequence of the encoded signal sequence (amino acid residues 1 to 22) and murine light chain (amino acid residues 23 to 235) is shown in SEQ ID NO: 58.

The murine Fd-CPG2 gene is prepared from the R6 variant of the CPG2 gene (d of Example 1) and the murine A5B7 Fd sequence in pAF1 (described in part d of Reference 25 Example 5 in International Patent Application WO 96/20011, Zeneca Limited). A PCR reaction with oligonucleotides SEQ ID NOS: 53 and 54 on pAF1 gives a 247 bp fragment. This is cut with HindIII and BamHI and cloned into similarly cut pUC19. The resulting plasmid is used to transform *E. coli* strain DH5α. The transformed cells are plated onto L agar plus ampicillin (100 μg/ml). A clone containing a plasmid with the correct sequence is named pUC19/muCH1/NcoI-AccIII(Fd). A second PCR with oligonucleotides SEQ ID NOS: 55 and 56 on pNG/VKss/CPG2/R6-neo (Example 1) gives a 265 bp fragment which is cut with HindIII and EcoRI and cloned into similarly cut pUC19 as above to give plasmid

pUC19/muCH1-linker-CPG2/AccIII-SacII. Plasmid pUC19/muCH1/Ncol-AccIII(Fd) is cut with HindIII and AccIII and the 258 bp fragment isolated by electrophoresis on a 1 % agarose gel. This fragment is cloned into HindIII and AccIII cut pUC19/muCH1-linker-CPG2/AccIII-SacII to give plasmid pUC19/muCH1-linker-CPG2/Ncol-SacII. A 956 bp fragment is isolated from pNG/VKss/CPG2/R6-neo by cutting it with SacII and EcoRI. This is cloned into SacII and EcoRI cut pUC19/muCH1-linker-CPG2/Ncol-SacII to give plasmid pUC19/muCH1-linker-RC/CPG2(R6). The complete gene construct is prepared by isolating a 498 bp HindIII to Ncol fragment from pAF1 and cloning it into HindIII and Ncol cut pUC19/muCH1-linker-RC/CPG2(R6). The resulting plasmid is used to transform *E. coli* strain DH5α. The transformed cells are plated onto L agar plus ampicillin (100 μg/ml). A clone containing a plasmid with the correct sequence and orientation is confirmed by DNA sequence analysis (SEQ ID NO: 59) and the plasmid named pUC19/muA5B7-RC/CPG2(R6).

The amino acid sequence of the encoded signal sequence (amino acid residues 1 to 19) and murine Fd-linker-CPG2 (amino acid residues 20 to 647) is shown in SEQ ID NO: 60.

15 Alternatively, the CPG2 gene sequence described in Example 1 can be obtained by total gene synthesis and converted to the R6 variant as described in d of Example 1. In this case, the

synthesis and converted to the R6 variant as described in d of Example 1. In this case, the base residue C at position 933 in SEQ ID NO: 59 is changed to G. The amino acid sequence of SEQ ID NO: 60 remains unaltered.

For expression in the pEE14 vector, the gene is first cloned into pEE6 (this is a derivative of pEE6.hCMV - Stephens and Cockett, 1989, Nucleic Acids Research 17, 7110, in which a HindIII site upstream of the hCMV promoter has been converted to a BglII site). Plasmid pUC19/muA5B7-RC/CPG2(R6) is cut with HindIII and EcoRI and the 1974 bp fragment isolated by electrophoresis on a 1 % agarose gel. This is cloned into HindIII and EcoRI cut pEE6 in *E. coli* strain DH5α to give plasmid pEE6/muA5B7-RC/CPG2(R6). The pEE14 co-expression vector is made by first cutting pEE6/muA5B7-RC/CPG2(R6) with BglII and BamHI and isolating the 4320 bp fragment on a 1 % agarose gel. This fragment is cloned into BglII and BamHI cut pEE14/A5B7muVkmuCK. The resulting plasmid is used to transform *E. coli* strain DH5α. The transformed cells are plated onto L agar plus ampicillin (100 μg/ml). A clone containing a plasmid with the correct sequence and orientation is confirmed by DNA sequence analysis and the plasmid named pEE14/muA5B7-RC/CPG2(R6).

For expression of (murine A5B7 Fab-CPG2)<sub>2</sub>, plasmid pEE14/muA5B7-RC/CPG2(R6) is used to transfect COS-7 or CHO cells as described in Example 48 of International Patent Application WO 97/42329, Zeneca Limited, published 13 November 1997. COS cell supernatants and CHO clone supernatants are assayed for activity as 5 described in Example 1 and shown to have CEA binding and CPG2 enzyme activity.

## Example 23

## Pharmaceutical composition

The following illustrate a representative pharmaceutical dosage form containing a 10 gene construct of the invention which may be used for therapy in combination with a suitable prodrug.

A sterile aqueous solution, for injection either parenterally or directly into tumour tissue, containing 107-1011 adenovirus particles comprising a gene construct as described in Example 1. After 3-7 days, three 1 g doses of prodrug are administered as sterile solutions at hourly intervals. Prodrug is selected from N-(4-[N,N-bis(2-iodoethyl)amino]-phenoxycarbonyl)-L-glutamic acid, N-(4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl)-L-glutamic-gamma-(3,5-dicarboxy)anilide or N-(4-[N,N-bis(2-chloroethyl)amino]-phenoxycarbonyl)-L-glutamic acid or a pharmaceutically acceptable salt thereof.

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#### SEQUENCE LISTING

```
(1) GENERAL INFORMATION:
  5
         (i) APPLICANT:
              (A) NAME: Zeneca Limited
               (B) STREET: 15 Stanhope Gate
              (C) CITY: London
 10
              (D) STATE: England
              (E) COUNTRY: United Kingdom
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              (G) TELEPHONE: 0171 304 5000
              (H) TELEFAX: 0171 304 5151
 15
              (I) TELEX: 0171 304 2042
        (ii) TITLE OF INVENTION: CHEMICAL COMPOUNDS
       (iii) NUMBER OF SEQUENCES: 60
 20
        (iv) COMPUTER READABLE FORM:
              (A) MEDIUM TYPE: Floppy disk
              (B) COMPUTER: IBM PC compatible
              (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 25
              (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
        (vi) PRIOR APPLICATION DATA:
             (A) APPLICATION NUMBER: GE 9709421.3
              (B) FILING DATE: 10-MAY-1997
 30
    (2) INFORMATION FOR SEQ ID NO: 1:
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              (D) TOPOLOGY: linear
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        (ii) MOLECULE TYPE: other nucleic acid
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        (i) SEQUENCE CHARACTERISTICS:
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		480
	TACGGCACCA TCACCGTGCT GTTCAACACC GACGAGGAAA AGGGTTCCTT CGGCTCGCGC	540

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10	GCCTACGC	CG C	GCTC	TCAG	G CA	AGCC	AGTG	ATC	GAGA	GCC	TGGG	CCTC	CC (	GGCT	TCGG	С	114	0
	TACCACAG									CGA	TTCC	GCGC	CG C	CTGT	ACAT	G	120	0
	GCTGCGCG	CC T	GATC	ATGG.	A TC	TGGG	CGCC	GGC	AAG								123	6
	(2) INFO	RMAT	ION	FOR .	SEQ	ID N	0: 1	0:										
15																		
	(i) SEQUENCE CHARACTERISTICS:																	
	(A) LENGTH: 412 amino acids																	
	(A) LENGTH: 412 amino acids (B) TYPE: amino acid																	
		(C	) ST	RAND	EDNE	ss:	sing	le										
20		(D	) TO	POLO	GY:	line	ar											
	U (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein																	
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	סא ם	: 10	:							
	Met	Asp	Phe	Gln	Val	Gln	Ile	Phe	Ser	Phe	Leu	Leu	Ile	Ser	Ala	Ser		
25	1				5					10					15	001		
	Val	Ile	Met	Ser	Arg	Gly	Gln	Lvs	Ara		Asn	Val	Len	Phe		A) =		
				20	_	•		-3-	25				200	30	J.II	nra		
	Ala	Thr	Asp	Glu	Gln	Pro	Ala	Val		Lvs	ፖስ r	î.eu	Glu		Lou	Va l		
			35					40		-,-		200	45	Dy 3	Deu	441		
30	Asn	Ile	Glu	Thr	Glv	Thr	Glv		Ala	Glu	Glv	Tle		λla	7.1.	C1		
		50			2		55			014	Gry	60	VIG	vra	та	GLY		
	Asn	Phe	Leu	Glu	Ala	Glu		1.178	Asn	Len	Gly		Th ~	Wal.	mL -	<b>N</b>		
	65					70	Deu	цуз	ASII	Ded	75	FIIG	Int	vai	Thr	_		
		Lvs	Ser	Ala	G) v		V=1	Wa 1	C)	7		<b>7</b> 1 -	1	<b>01</b>	_	80		
35		-,0	001		85	Deu	Vai	AGI	GLY		ASI	TTE	vaı	GIÀ		He		
	Lvs	Glv	Ara	Glv		1	A on	T 0	1	90	14	•			95			
	-,5	OLy	ary	Gly 100	GIY	Lys	ASII	ren		Leu	Met	Ser	His		Asp	Thr		
	(Let)	T	Tou		<b>63</b>	<b>7</b> 1 -			105		_			110				
	٧٨١	1 1 1	115	Lys	GTA	me	reu		ьуз	Ala	Pro	Phe		Val	Glu	Gly		
40	Acn	7.110		m	<b>61</b>	D	۵.	120					125					
70	ASP		ATA	Tyr.	GIÀ	Pro		Ile	Ala	Asp	Asp	Lys	Gly	Gly	Asn	Ala		
	**- 1	130	_				135					140						
		тте	Leu	His	Thr		Lys	Leu	Leu	Lys	Glu	Tyr	Gly	Val	Arg	Asp		
	145					150					155					160		
15	Tyr	Gly	Thr	Ile		Val	Leu	Phe	Asn	Thr	Asp	Glu	Glu	Lys	Gly	Ser		
45					165					170					175			
	Phe	Gly	Ser	Arg	Asp	Leu	Ile	Gln	Glu	Glu	Ala	Lys	Leu	Ala	Asp	Tyr		
				180					185					190				

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		Val	Leu	Ser 195	Phe	Glu	Pro	Thr		Ala	Gly	Asp	Glu		Leu	Ser	Leu
		G) u	Th-			T) -	21-		200					205			
		Gry	210	ser	Gly	116	Ala		vaı	Gin	Val	GIn		Thr	Gly	Lys	Ala
5		C		71-	<b>61</b>			215					220				
J			113	ATA	Gly	Ala		Pro	Glu	Leu	Gly		Asn	Ala	Leu	Val	Glu
		225			_		230					235					240
		Ala	Ser	Asp	Leu		Leu	Arg	Thr	Met	Asn	Ile	Asp	qeA	Lys	Ala	Lys
			_			245					250					255	
10		Asn	Leu	Arg	Phe	Gln	Trp	Thr	Ile	Ala	Lys	Ala	Gly	Gln	Val	Ser	Asn
10					260					265					270		
		Ile	Ile	Pro	Ala	Ser	Ala	Thr	Leu	Asn	Ala	Asp	Val	Arg	Tyr	Ala	Arg
				275					280					285			
		Asn		Asp	Phe	Asp	Ala	Ala	Met	Lys	Thr	Leu	Glu	Glu	Arg	Ala	Gln
			290					295					300				
15		Gln	Lys	Lys	Leu	Pro	Glu	Ala	Asp	Val	Lys	Val	Ile	Val	Thr	Arg	Gly
		305					310					315					320
		Arg	Pro	Ala	Phe	Asn	Ala	Gly	Glu	Gly	Gly	Lys	Lys	Leu	Val	Asp	Lys
						325					330					335	
		Ala	Val	Ala	Tyr	Tyr	Lys	Glu	Ala	Gly	Gly	Thr	Leu	Gly	Val	Glu	Glu
20					340					345					350		
		Arg	Thr	Gly	Gly	Gly	Thr	Asp	Ala	Ala	Tyr	Ala	Ala	Leu	Ser	Gly	Lys
				355					360					365		-	•
		Pro	Val	Ile	Glu	Ser	Leu	Gly	Leu	Pro	Gly	Phe	Gly	Tyr	His	Ser	Asp
			370					375					380	-			
25		Lys	Ala	Glu	Tyr	Val	Asp	Ile	Ser	Ala	Ile	Pro	Arg	Ara	Leu	Tvr	Met
		385					390					395	•	•		- 3 -	400
		Ala	Ala	Arg	Leu	Ile	Met	Asp	Leu	Gly	Ala	Glv	Lvs				
						405		-		-	410	-	-,				
30	(2)	INFOR	MATI	ON F	OR S	EQ I	D NO	: 11	. :								
					СНА												
			(A)	LEN	IGTH:	21	base	pai	rs								
					E: n												
			(C)	STR	ANDE	DNES	S: s	ingl	e								
35					OLOG					•							
	(	(ii)			TYP				leic	aci	d						
					DES							•					
	CCACI	CTCA	C AG	TGAG	CTCG	G											

21

40

- (2) INFORMATION FOR SEQ ID NO: 12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 55 base pairs
    - (B) TYPE: nucleic acid
- 45 (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12: ACCGCTACCG CCACCACCAG AGCCACCACC GCCAACTGTC TTGTCCACCT TGGTG 55 5 (2) INFORMATION FOR SEQ ID NO: 13: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13: ACCCCCTCTA GAGTCGAC 18 15 (2) INFORMATION FOR SEQ ID NO: 14: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 54 base pairs (B) TYPE: nucleic acid 20 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14: 25 TCTGGTGGTG GCGGTAGCGG TGGCGGGGGT TCCCAGAAGC GCGACAACGT GCTG 54 (2) INFORMATION FOR SEQ ID NO: 15: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1929 base pairs 30 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: other nucleic acid (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15: 35 ATGGAGTTGT GGCTGAACTG GATTTTCCTT GTAACACTTT TAAATGGTAT CCAGTGTGAG GTGAAGCTGG TGGAGTCTGG AGGAGGCTTG GTACAGCCTG GGGGTTCTCT GAGACTCTCC TGTGCAACTT CTGGGTTCAC CTTCACTGAT TACTACATGA ACTGGGTCCG CCAGCCTCCA 180 GGAAAGGCAC TTGAGTGGTT GGGTTTTATT GGAAACAAAG CTAATGGTTA CACAACAGAG 240 40 tacagtgcat ctgtgaaggg tcggttcacc atctccagag ataaatccca aagcatcctc TATCTTCAAA TGAACACCCT GAGAGCTGAG GACAGTGCCA CTTATTACTG TACAAGAGAT 360 AGGGGGCTAC GGTTCTACTT TGACTACTGG GGCCAAGGCA CCACTCTCAC AGTGAGCTCG 420 GCTAGCACCA AGGGACCATC GGTCTTCCCC CTGGCCCCCT GCTCCAGGAG CACCTCCGAG 480 AGCACAGCCG CCCTGGGCTG CCTGGTCAAG GACTACTTCC CCGAACCGGT GACGGTGTCG 45 TGGAACTCAG GCGCTCTGAC CAGCGGCGTG CACACCTTCC CGGCTGTCCT ACAGTCCTCA 600 GGACTCTACT CCCTCAGCAG CGTCGTGACG GTGCCCTCCA GCAACTTCGG CACCCAGACC 660

TACACCTGCA ACGTAGATCA CAAGCCCAGC AACACCAAGG TGGACAAGAC AGTTGGCGGT

720

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	GGTGGCTCTG	GTGGTGGCGG	TAGCGGTGGC	GGGGGTTCCC	AGAAGCGCGA	CAACGTGCTG	780
	TTCCAGGCAG	CTACCGACGA	GCAGCCGGCC	GTGATCAAGA	CGCTGGAGAA	GCTGGTCAAC	840
	ATCGAGACCG	GCACCGGTGA	CGCCGAGGGC	ATCGCCGCTG	CGGGCAACTT	CCTCGAGGCC	900
	GAGCTCAAGA	ACCTCGGCTT	CACGGTCACG	CGAAGCAAGT	CGGCCGGCCT	GGTGGTGGGC	960
5	GACAACATCG	TGGGCAAGAT	CAAGGCCGC	GGCGGCAAGA	ACCTGCTGCT	GATGTCGCAC	1020
	ATGGACACCG	TCTACCTCAA	GGGCATTCTC	GCGAAGGCCC	CGTTCCGCGT	CGAAGGCGAC	1080
					ACGCGGTCAT		1140
					CCATCACCGT		1200
					TCCAGGAAGA		1260
10	GCCGACTACG						1320
					AGGCCTCGCA		1380
					ACCTCGTGCT		1440
					CCATCGCCAA		1500
					ACGTGCGCTA		1560
15	GAGGACTTCG						1620
					CCTTCAATGC		1680
					AAGCCGGCGG		1740
					CCGCGCTCTC		1800
					GCGACAAGGC		1860
20					GCCTGATCAT		1920
	GCCGGCAAG				_		1929

## (2) INFORMATION FOR SEQ ID NO: 16:

(i) SEQUENCE CHARACTERISTICS:

25

(A) LENGTH: 643 amino acids

(B) TYPE: amino acid ·

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

Met Glu Leu Trp Leu Asn Trp Ile Phe Leu Val Thr Leu Leu Asn Gly 10 Ile Gln Cys Glu Val Lys Leu Val Glu Ser Gly Gly Gly Leu Val Gln 35 25 Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Thr Ser Gly Phe Thr Phe 40 Thr Asp Tyr Tyr Met Asn Trp Val Arg Gln Pro Pro Gly Lys Ala Leu 55 40 Glu Trp Leu Gly Phe Ile Gly Asn Lys Ala Asn Gly Tyr Thr Thr Glu 70 75 Tyr Ser Ala Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Lys Ser 85 90 Gin Ser Ile Leu Tyr Leu Gin Met Asn Thr Leu Arg Ala Glu Asp Ser 45 105 Ala Thr Tyr Tyr Cys Thr Arg Asp Arg Gly Leu Arg Phe Tyr Phe Asp 120

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	Tyr			Gln	Gly	Thi	Thr		Thr	Val	Ser	Ser	Ala	Ser	Thi	Ly
	Glu	130			n		135					140				
	145		, ser	val	Pne	150	Leu	Ala	Pro	Cys			Ser	Thr	Ser	
5			· Ala	Δla	Leu			T 011	17.01	*	155		_,			160
•	561	••••	, AIG	. AIG	165		' Cys	Leu	. vaı	ьуя 170		Tyr	Phe	Pro		
	Val	Thr	Val	Ser			Ser	Glv	- D 1-a				<b>.</b>	. 17-1	175	
				180				Oly	185		1111	261	GT	7 vai 190		Thi
	Phe	Pro	Ala			Gln	Ser	Ser			TVY	Ser	ī.e.			. (7-1
10			195					200			.,_	501	205		Set	. va.
	Val	Thr	Val	Pro	Ser	Ser	Asn	Phe	Gly	Thr	Gln	Thr			Cvs	Asr
		210					215					220			-3-	
	Val	Asp	His	Lys	Pro	Ser	Asn	Thr	Lys	Val	Asp	Lys	Thr	Val	Gly	Gly
	225					230					235					240
15	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gln	Lys	Arg
					245					250					255	
	Asp	Asn	Val			Gln	Ala	Ala	Thr	qzA	Glu	Gln	Pro	Ala	Val	Ile
	r	-		260		_			265					270		
20	гÀг	Thr	275		Lys	Leu	Val		Ile	Glu	Thr	Gly		Gly	Asp	Ala
20	Glu	Glu			21-	<b>31</b> -	G)	280		_			285			
	010	290		nt a	MIG	WIG	Gly 295	Asn	Pne	ren	GIu	•	Glu	Leu	Lys	Asn
	Leu			Thr	Val	Thr	Arg	Ser	Luc	Sar	A 1 =	300	7 0	17-1	17-1	<b>01</b>
	305	•				310		Jer	Lys	Set	315	GIĀ	nen	Val	vaı	320
25	Asp	Asn	Ile	Val	Gly	•	Ile	Lvs	Glv	Ara		Glv	Lvs	Asn	ī.eu	
					325	_		•	•	330	1	1	-,-		335	
•	Leu	Met	Ser	His	Met	Asp	Thr	Val	Tyr	Leu	Lys	Gly	Ile	Leu		
				340					345					350		_
••	Ala	Pro	Phe	Arg	Val	Glu	Gly	Asp	Lys	Ala	Tyr	Gly	Pro	Gly	Ile	Ala
30			355					360					365			
	Asp		Lys	Gly	Gly	Asn	Ala	Val	Ile	Leu	His	Thr	Leu	Lys	Leu	Leu
	•	370	_				375					380				
		Glu	Tyr	Gly	Val		Asp	Туг	Gly	Thr		Thr	Val	Leu	Phe	Asn
35	385 The	Acn	C1	C1	*	390	0			_	395	_				400
,,	****	изр	GIU	GIU	405	GIY	Ser	Pne	GIY		Arg	Asp	Leu	Ile		
	Glu	Ala	Lvs	Leu		Asn	Tyr	Val	Lou	410	Dho	C1	D	mh	415	
			-,-	420		wob	. y .	AGI	425	ser	rne	GIU	PIO	430	ser	Ala
	Gly	Asp	Glu		Leu	Ser	Leu	Glv		Ser	Glv	Tle	Δla		Va 1	Gln
40			435	_				440			,		445	- 7 -	,	01
	Val	Gln	Ile	Thr	Gly	Lys	Ala	Ser	His	Ala	Gly	Ala		Pro	Glu	Leu
		450					455				-	460				
	Gly	Val	Asn	Ala	Leu	Val	Glu	Ala	Ser	Asp	Leu	Val	Leu	Arg	Thr	Met
	465					470					475					480
45	Asn	Ile	Asp	qeA	Lys	Ala	Lys	Asn	Leu	Arg	Phe	Gln	Trp	Thr	Ile	Ala
					485					490					495	
	Lys	Ala	Glv	Gln	Val	Ser	Asn	Tle	Tla	Pro	בומ	Sar	λla	Th-	T 011	7.00

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				500					505					510			
	Ala	Asp	Val	Arg	Tyr	Ala	Ara	Asn		Asp	Phe	Asn	Ala		Mot	T	
			515	_	-		5	520		· · · · ·		nap	525	ΛIα	nec	LyS	
	Thr	Leu	Glu	Glu	Arg	Ala	Gln	Gln	Lvs	Lvs	Leu	Pro		Δla	Asn	Val	
5		530			_		535		-,-	-,-		540	CIG	AIG	vab	val	
	Lys	Val	Ile	Val	Thr	Arg		Ara	Pro	Ala	Phe		Δla	Gly	Glu	C1	
	545					550					555			Gry	GIU	560	
	Gly	Lys	Lys	Leu	Val	Asp	Lys	Ala	Val	Ala		Tvr	I.vs	Glu	Δ1 a		
					565	-	•			570	-1-	.,.	-,,	014	575	GIŞ	
10	Gly	Thr	Leu	Gly	Val	Glu	Glu	Arg	Thr		Glv	Glv	Thr	Asp		Δla	
				580				-	585	•		2		590		niu	
	Tyr	Ala	Ala	Leu	Ser	Gly	Lys	Pro	Val	Ile	Glu	Ser	Leu		Leu	Pro	
			595					600					605	,	204		
	Gly	Phe	Gly	Tyr	His	Ser	Asp	Lys	Ala	Glu	Tyr	Val	Asp	Ile	Ser	Ala	
15		610	•				615				-	620	•				
	Ile	Pro	Arg	Arg	Leu	Tyr	Met	Ala	Ala	Arg	Leu	Ile	Met	Asp	Leu	Glv	
	625					630					635			•		640	
	Ala	Gly	Lys														
•																	
20				•													
	(2) INFOR																
	(i)					ERIS											
						bas		irs									
25						ic a											
25						S: s	_	.e									
	1221					inea	_										
	(ii)																
	(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	17:							
30	ATGGATTTT	מג חי	GTGC	ימכטמי	ጥጥጥ	ראכר	TTTC	CTCC	m > > m	<b></b> .							
	AGAGGACAA	A CT	GTTC	TCTC	CCA	CTCT	CCD	CIGO	TAMI	CT C	TGCT	TCAG	T CA	TAAT	GTCC		60
	GTCACAATG	A CT	TGCA	GGGC	CAG	CTCA	AGT	GCAA	ርጥጥ ነርርነ	ים כא יים	TOOM	TC1C	C AG	BGGA BCCB	GAAG		120
	CCAGGTTCC	T CC	CCCA	AATC	CTG	GATT	TAT	GCCA	САТС	CA A	CCTG	. ദേശ്വ വേസത	ת ככ כידכ	CACT	CCCT		180
	GCTCGCTTC																300
35	GCTGAAGAT	G CT	GCCA	CTTA	TTA	CTGC	CAA	CATT	GGAG	TA G	TAAA	מבפת	ר המ	CCTT	CCCT		360
	GGAGGCACC	A AG	CTCG	AGAT	CAA	ACGG.	ACT	GTGG	CTGC	AC C	ATCT	STCT	T CA	TOTT	ccce		420
	CCATCTGAT	G AG	CAGT	TGAA	ATC	TGGA	ACT	GCCT	CTGT	TG T	GTGC	CTGC'	T GA	ATAA	CTTC		480
	TATCCCAGA																540
	CAGGAGAGT																600
40	ACGCTGAGC	A AA	GCAG	ACTA	CGA	GAAA	CAC .	AAAG'	rcta	CG C	CTGC	GAAG'	r ca	CCCA	rcag		660
	GGCCTGAGT	T CG	CCCG	TCAC	AAA	GAGC'	TTC .	AACA	GGGG.	AG A	GTGT						705

## (2) INFORMATION FOR SEQ ID NO: 18:

- (i) SEQUENCE CHARACTERISTICS:
- 45 (A) LENGTH: 235 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single

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(D) TOPOLOGY: linear
        (ii) MOLECULE TYPE: protein
        (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:
 5
        Met Asp Phe Gln Val Gln Ile Phe Ser Phe Leu Leu Ile Ser Ala Ser
                                           10
        Val Ile Met Ser Arg Gly Gln Thr Val Leu Ser Gln Ser Pro Ala Ile
                                       25
        Leu Ser Ala Ser Pro Gly Glu Lys Val Thr Met Thr Cys Arg Ala Ser
10
                                   40
        Ser Ser Val Thr Tyr Ile His Trp Tyr Gln Gln Lys Pro Gly Ser Ser
                               55
        Pro Lys Ser Trp Ile Tyr Ala Thr Ser Asn Leu Ala Ser Gly Val Pro
                          70
                                            75
15
        Ala Arg Phe Ser Gly Ser Gly Ser Gly Thr Ser Tyr Ser Leu Thr Ile
                      85 90
        Ser Arg Val Glu Ala Glu Asp Ala Ala Thr Tyr Tyr Cys Gln His Trp
                              105
        Ser Ser Lys Pro Pro Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys
20
                                   120
        Arg Thr Val Ala Ala Pro Ser Val Phe Ile Phe Pro Pro Ser Asp Glu
                               135
        Gln Leu Lys Ser Gly Thr Ala Ser Val Val Cys Leu Leu Asn Asn Phe
                          150
                                             155
25
        Tyr Pro Arg Glu Ala Lys Val Gln Trp Lys Val Asp Asn Ala Leu Gln
                       165
                                         170
        Ser Gly Asn Ser Gln Glu Ser Val Thr Glu Gln Asp Ser Lys Asp Ser
                                       185
        Thr Tyr Ser Leu Ser Ser Thr Leu Thr Leu Ser Lys Ala Asp Tyr Glu
30
                                   200
        Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser Ser
                              215
        Pro Val Thr Lys Ser Phe Asn Arg Gly Glu Cys
                           230
35
   (2) INFORMATION FOR SEQ ID NO: 19:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 39 base pairs
            (B) TYPE: nucleic acid
40
            (C) STRANDEDNESS: single
            (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: other nucleic acid
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:
```

39

45 AAGCTTGAAT TCGCCGCCAC TATGGATTTT CAAGTGCAG

(2) INFORMATION FOR SEQ ID NO: 20:

	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 44 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
5	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi: SEQUENCE DESCRIPTION: SEQ ID NO: 20:	
	TTAATTGGAT CCGAGCTCCT ATTAACACTC TCCCCTGTTG AAGC	44
10	·	
	(2) INFORMATION FOR SEQ ID NO: 21:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 50 base pairs	
	(B) TYPE: nucleic acid	
15	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:	
20	AAGCTTCCGG ATCCCTGCAG CCATGGAGTT GTGGCTGAAC TGGATTTTCC	50
	(2) INFORMATION FOR SEQ ID NO: 22:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 38 base pairs	
25	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
•	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:	
30		
	AAGCTTAGTC TAGATTATCA CTTGCCGGCG CCCAGATC	38
	(2) INFORMATION FOR SEQ ID NO: 23:	
٠.	(i) SEQUENCE CHARACTERISTICS:	
35	(A) LENGTH: 46 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: other nucleic acid	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:	
	CCCCCATCC ACAMCMCACC MCCMCMACAC CMCCACACAC	,
	CGGGGGATCC AGATCTGAGC TCCTGTAGAC GTCGACATTA ATTCCG	46
	(2) INFORMATION FOR SEQ ID NO: 24:	
15	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 30 base pairs	

(B) TYPE: nucleic acid

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(C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

5

### GGAAAATCCA GTTCAGCCAC AACTCCATGG

30

60

### (2) INFORMATION FOR SEQ ID NO: 25:

(i) SEQUENCE CHARACTERISTICS:

10

- (A) LENGTH: 1926 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid
- 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

ATGAAGTTGT GGCTGAACTG GATTTTCCTT GTAACACTTT TAAATGGAAT TCAGTGTGAG GTGCAGCTGC AGCAGTCTGG GGCAGAGCTT GTGAGGTCAG GGGCCTCAGT CAAGTTGTCC 120 TGCACAGCTT CTGGCTTCAA CATTAAAGAC AACTATATGC ACTGGGTGAA GCAGAGGCCT 180 20 gaacagggcc tggagtggat tgcatggatt gatcctgaga atggtgatac tgaatatgcc 240 CCGAAGTTCC GGGGCAAGGC CACTTTGACT GCAGACTCAT CCTCCAACAC AGCCTACCTG 300 CACCTCAGCA GCCTGACATC TGAGGACACT GCCGTCTATT ACTGTCATGT CCTGATCTAT 360 GCTGGTTATT TGGCTATGGA CTACTGGGGT CAAGGAACCT CAGTCGCCGT GAGCTCGGCT 420 AGCACCAAGG GACCATCGGT CTTCCCCCTG GCCCCCTGCT CCAGGAGCAC CTCCGAGAGC 480 25 ACAGCCGCCC TGGGCTGCCT GGTCAAGGAC TACTTCCCCG AACCGGTGAC GGTGTCGTGG AACTCAGGCG CTCTGACCAG CGGCGTGCAC ACCTTCCCGG CTGTCCTACA GTCCTCAGGA 600 CTCTACTCCC TCAGCAGCGT CGTGACGGTG CCCTCCAGCA ACTTCGGCAC CCAGACCTAC 660 ACCTGCAACG TAGATCACAA GCCCAGCAAC ACCAAGGTGG ACAAGACAGT TGGCGGTGGT 720 GGCTCTGGTG GTGGCGGTAG CGGTGGCGGG GGTTCCCAGA AGCGCGACAA CGTGCTGTTC 780 30 CAGGCAGCTA CCGACGAGCA GCCGGCCGTG ATCAAGACGC TGGAGAAGCT GGTCAACATC 840 GAGACCGGCA CCGGTGACGC CGAGGGCATC GCCGCTGCGG GCAACTTCCT CGAGGCCGAG 900 CTCAAGAACC TCGGCTTCAC GGTCACGCGA AGCAAGTCGG CCGGCCTGGT GGTGGGCGAC 960 AACATCGTGG GCAAGATCAA GGGCCGCGGC GGCAAGAACC TGCTGCTGAT GTCGCACATG 1020 GACACCGTCT ACCTCAAGGG CATTCTCGCG AAGGCCCCGT TCCGCGTCGA AGGCGACAAG 1080 35 GCCTACGGCC CGGGCATCGC CGACGACAAG GGCCGCAACG CGGTCATCCT GCACACGCTC AAGCTGCTGA AGGAATACGG CGTGCGCGAC TACGGCACCA TCACCGTGCT GTTCAACACC 1200 GACGAGGAAA AGGGTTCCTT CGGCTCGCGC GACCTGATCC AGGAAGAAGC CAAGCTGGCC 1260 GACTACGTGC TCTCCTTCGA GCCCACCAGC GCAGGCGACG AAAAACTCTC GCTGGGCACC 1320 TCGGGCATCG CCTACGTGCA GGTCCAGATC ACCGGCAAGG CCTCGCATGC CGGCGCCGCG 1380 40 CCCGAGCTGG GCGTGAACGC GCTGGTCGAG GCTTCCGACC TCGTGCTGCG CACGATGAAC 1440 ATCGACGACA AGGCGAAGAA CCTGCGCTTC CAGTGGACCA TCGCCAAGGC CGGCCAGGTC 1500 TCGAACATCA TCCCCGCCAG CGCCACGCTG AACGCCGACG TGCGCTACGC GCGCAACGAG 1560 GACTTCGACG CCGCCATGAA GACGCTGGAA GAGCGCGCGC AGCAGAAGAA GCTGCCCGAG 1620 GCCGACGTGA AGGTGATCGT CACGCGCGGCC CGCCCGGCCT TCAATGCCGG CGAAGGCGGC 1680

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										- /-	-					
	AAGAAGCT	GG I	CGAC	AAGC	C GG	TGGC	CTAC	TAC	AAGG	AAG	CCGC	CGGC	AC C	CTG	GCG	r <b>G</b>
	GAAGAGCG															
	ATCGAGAG															
5	ATCAGCGC GGCAAG	.011 1	1000	CGCC	.0 ((	IGIA	CAIG	GCI	GCGC	GCC	TGAT	CATG	iGA T	CTGC	GCGC	:C
,	GGCAAG															
	(2) INFO	RMAT	ION	FOR	SEQ	ID N	0: 2	6:								
	(i)	SEQ	UENC	E CH	ARAC	TERI	STIC	s:								
		(A	) LE	NGTH	: 64	2 am	ino	acid	s							
10			) TY													
		(C	) ST	RAND	EDNE	SS:	sing	le								
		(D	) TO	POLO	GY:	line	ar									
			ECUL													
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 26	:					
15																
	Met	Lys	Leu	Trp	Leu	Asn	Trp	Ile	Phe	Leu	Val	Thr	Leu	Leu	Asn	Gly
	1				5					10					15	
	Ile	Gln	Cys	Glu	Val	Gln	Leu	Gln	Gln	Ser	Gly	Ala	Glu	Leu	Val	Arg
				20					25					30		
20	Ser	Gly	Ala	Ser	Val	Lys	Leu	Ser	Cys	Thr	Ala	Ser	Gly	Phe	Asn	Ile
			35					40					45			
	Lys	Asp	Asn	Tyr	Met	His	Trp	Val	Lys	Gln	Arg	Pro	Glu	Gln	Gly	Leu
		50					55					60				
	Glu	Trp	Ile	Ala	Trp	Ile	Asp	Pro	Glu	Asn	Gly	Asp	Thr	Glu	Tyr	Ala
25	65					70					75					80
	Pro	Lys	Phe	Arg	Gly	Lys	Ala	Thr	Leu	Thr	Ala	Asp	Ser	Ser	Ser	Asn
					85					90					95	
	Thr	Ala	Tyr		His	Leu	Ser	Ser	Leu	Thr	Ser	Glu	Asp	Thr	Ala	Val
20	_			100					105					110		
30	Tyr	Tyr	Cys	His	Val	Leu	Ile	Tyr	Ala	Gly	Tyr	Leu	Ala	Met	Asp	Tyr
	_		115					120					125			
	Trp		Gln	Gly	Thr	Ser	Val	Ala	Val	Ser	Ser	Ala	Ser	Thr	Lys	Gly
	_	130					135					140				
25		Ser	Val	Phe	Pro	Leu	Ala	Pro	Суз	Ser	Arg	Ser	Thr	Ser	Glu	Ser
35	145					150					155					160
	Thr	Ala	Ala	Leu		Суз	Leu	Val	Lys		Tyr	Phe	Pro	Glu	Pro	Val
					165					170					175	
	Thr	Val	Ser		Asn	Ser	Gly	Ala	Leu	Thr	Ser	Gly	Val	His	Thr	Phe
40				180					185					190		
40	Pro	Ala	Val	Leu	Gln	Ser	Ser		Leu	Tyr	Ser	Leu	Ser	Ser	Val	Val
			195	•				200					205			
	Thr		Pro	Ser	Ser	Asn	Phe	Gly	Thr	Gln	Thr	Tyr	Thr	Cys	Asn	Val
		210					215					220				
15		His	Lys	Pro	Ser		Thr	Lys	Val	qzA	l,ys	Thr	Val	Gly	Gly	Gly
45	225	_				230					235					240
	Gly	Ser	Gly	Gly		Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gln	Lys	Arg	Asp
					245					250					255	

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	Asn	Val	Leu			Ala	Ala	Thr			Gln	Pro	Ala	Val	Ile	Lys
	mb	•	۵.	260					265					270		
	inr	reu			Leu	Val	Asn			Thr	Gly	Thr	Gly	Asp	Ala	Glu
5	C1	T1.	275				_	280					285			
,	GIY			Ala	Ala	Gly			Leu	Glu				Lys	Asn	Leu
	<b>6</b> 1	290					295					300				
		rne	Thr	Val	Thr			Lys	Ser	Ala			Val	Val	Gly	Asp
	305	<b>T</b> 1-	**- 1	۵.		310					315					320
10	Asn	ile	vaı	GIY		Ile	Lys	Gly	Arg		Gly	Lys	Asn	Leu	Leu	Leu
10	Mak	0	<b>11</b> 2 -		325			_		330					335	
	riet	Ser	піз		Asp	Thr	Val	Tyr			Gly	Ile	Leu			Ala
`	Pro	Pho	7~~	340	C1	G3		•	345			_		350		
	110	rne	355	var	GIU	GIĀ	Asp		Ala	Tyr	Gly	Pro			Ala	Asp
15	Δen	Tue		C1	7.00	77-		360	•			_	365			
••	nsp	370	GIY	GIY	ASII	Ala	375	ite	Leu	HIS	Thr			Leu	Leu	Lys
	Glu		Gly	V = 1	7			C1	m	T1-	<b>m</b> >	380				
	385		OLY	441	ALY	390		GIY	Int	TIG		Val	Leu	Phe	Asn	Thr
		Glu	Glu	Lve	Glv			G) v	80=	N == ~	395	T	<b>7</b> 1 -	<b>01</b>		400 Glu
20				270	405	561	riie	GIY	ser	410	АЗР	ren	iie	GIN		Glu
	Ala	Lvs	Leu	Ala		Tur	Val	T.em	Sar		C) is	Dwa	mh -	c	415	Gly
		-,-		420		- , -	• • • •	Deu	425	rine	Gru	FIG	Int	3er	Ala	GIÀ
	qzA	Glu	Lvs		Ser	Leu	Glv	Thr		Glv	Tla	A1 =	T		C1-	Val
	=		435				,	440		013	110	niu	445	Val	GIII	val
25	Gln	Ile	Thr	Gly	Lys	Ala	Ser		Ala	Glv	λla	Ala		Glu	Len	Gly
		450		-	-		455			,		460	110	014	beu	GIY
	Val	Asn	Ala	Leu	Val	Glu	Ala	Ser	Asp	Leu	Val		Ara	Thr	Met	Asn
	465					470			•		475		3			480
	Ile	Asp	Asp	Lys	Ala	Lys	Asn	Leu	Arg	Phe	Gln	Trp	Thr	Ile	Ala	
30					485				-	490		•			495	-10
	Ala	Gly	Gln	Val	Ser	Asn	Ile	Ile	Pro	Ala	Ser	Ala	Thr	Leu	Asn	Ala
				500					505					510		
	Asp	Val	Arg	Tyr	Ala	Arg	Asn	Glu	Asp	Phe	Asp	Ala	Ala	Met	Lys	Thr
			515					520		•			525		_	
35	Leu	Glu	Glu	Arg	Ala	Gln	Gln	Lys	Lys	Leu	Pro	Glu	Ala	Asp	Val	Lys
		530					535					540				
	Val	Ile	Val	Thr	Arg	Gly	Arg	Pro	Ala	Phe	Asn	Ala	Gly	Glu	Gly	Gly
•	545					550					555					560
	Lys	Lys	Leu	Val	Asp	Lys	Ala	Val	Ala	Tyr	Tyr	Lys	Glu	Ala	Gly	Gly
40					565					570					575	
	Thr	Leu	Gly	Val	Glu	Glu	Arg	Thr	Gly	Gly	Gly	Thr	Asp	Ala	Ala	Tyr
				580					585				•	590		
	Ala	Ala		Ser	Gly	Lys	Pro	Val	Ile	Glu	Ser	Leu	Gly	Leu	Pro	Gly
15			595					600					605			
45	Phe		Tyr	His	Ser	Asp		Ala	Glu	Tyr	Val	Asp	Ile	Ser	Ala	Ile
		610					615					620				
	Pro	Arg	Arg	Leu	Tyr	Met	Ala	Ala	Arg	Leu	Ile	Met	Asp	Leu	Gly	Ala

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625 630 635 640 Gly Lys

5 (2) INFORMATION FOR SEQ ID NO: 27:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 39 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

10 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

AAGCTTGGAA TTCAGTGTCA GGTCCAACTG CAGCAGCCT

39

(2) INFORMATION FOR SEQ ID NO: 28:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 54 base pairs

(B) TYPE: nucleic acid

20 (C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

25 GCTACCGCCA CCTCCGGAGC CACCACCGCC CCGTTTGATC TCGAGCTTGG TGCC

54

(2) INFORMATION FOR SEQ ID NO: 29:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 58 base pairs

30

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

35

TCCGGAGGTG GCGCTAGCGG TGGCGGGGGT TCCCAGAAGC GCGACAACGT GCTGTTCC

58

(2) INFORMATION FOR SEQ ID NO: 30:

(i) SEQUENCE CHARACTERISTICS:

40

(A) LENGTH: 24 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

CCTCGAGGAA TTCTTTCACT TGCC

24

(2) INFORMATION FOR SEQ ID NO: 31:

```
(i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 2019 base pairs
   5
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: other nucleic acid
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:
 10
    ATGAAGTTGT GGCTGAACTG GATTTTCCTT GTAACACTTT TAAATGGAAT TCAGTGTCAG
    GTCCAACTGC AGCAGCCTGG GGCTGAACTG GTGAAGCCTG GGGCTTCAGT GCAGCTGTCC
                                                                           120
    TGCAAGGCTT CTGGCTACAC CTTCACCGGC TACTGGATAC ACTGGGTGAA GCAGAGGCCT
                                                                           180
    GGACAAGGCC TTGAGTGGAT TGGAGAGGTT AATCCTAGTA CCGGTCGTTC TGACTACAAT
                                                                           240
15 GAGAAGTTCA AGAACAAGGC CACACTGACT GTAGACAAAT CCTCCACCAC AGCCTACATG
                                                                           300
    CAACTCAGCA GCCTGACATC TGAGGACTCT GCGGTCTATT ACTGTGCAAG AGAGAGGGCC
                                                                           360
    TATGGTTACG ACGATGCTAT GGACTACTGG GGCCAAGGGA CCACGGTCAC CGTCTCCTCA
                                                                           420
    GGTGGCGGTG GCTCGGGCGG TGGTGGGTCG GGTGGCGGCG GATCTGACAT TGAGCTCTCA
                                                                           480
    CAGTCTCCAT CCTCCCTGGC TGTGTCAGCA GGAGAGAGG TCACCATGAG CTGCAAATCC
                                                                           540
 20 agtcagagtc tectcaacag tagaaceega aagaactaet tegettegta eeagcagaga
                                                                           600
    CCAGGGCAGT CTCCTAAACT GCTGATCTAT TGGGCATCCA CTAGGACATC TGGGGTCCCT
                                                                           660
    GATCGCTTCA CAGGCAGTGG ATCTGGGACA GATTTCACTC TCACCATCAG CAGTGTGCAG
                                                                          720
    GCTGAAGACC TGGCAATTTA TTACTGCAAG CAATCTTATA CTCTTCGGAC GTTCGGTGGA
                                                                          780
    GGCACCAAGC TCGAGATCAA ACGGGGCGGT GGTGGCTCCG GAGGTGGCGG TAGCGGTGGC
                                                                          840
 25 GGGGGTTCCC AGAAGCGCGA CAACGTGCTG TTCCAGGCAG CTACCGACGA GCAGCCGGCC
                                                                          900
    GTGATCAAGA CGCTGGAGAA GCTGGTCAAC ATCGAGACCG GCACCGGTGA CGCCGAGGGC
                                                                          960
    ATCGCCGCTG CGGGCAACTT CCTCGAGGCC GAGCTCAAGA ACCTCGGCTT CACGGTCACG
                                                                         1020
    CGAAGCAAGT CGGCCGGCCT GGTGGTGGGC GACAACATCG TGGGCAAGAT CAAGGGCCGC
                                                                         1080
    GGCGGCAAGA ACCTGCTGCT GATGTCGCAC ATGGACACCG TCTACCTCAA GGGCATTCTC
                                                                         1140
 30 GCGAAGGCCC CGTTCCGCGT CGAAGGCGAC AAGGCCTACG GCCCGGGCAT CGCCGACGAC
                                                                         1200
    AAGGGCGGCA ACGCGGTCAT CCTGCACACG CTCAAGCTGC TGAAGGAATA CGGCGTGCGC
                                                                         1260
    GACTACGGCA CCATCACCGT GCTGTTCAAC ACCGACGAGG AAAAGGGTTC CTTCGGCTCG
                                                                         1320
    CGCGACCTGA TCCAGGAAGA AGCCAAGCTG GCCGACTACG TGCTCTCCTT CGAGCCCACC
                                                                         1380
    AGCGCAGGCG ACGAAAAACT CTCGCTGGGC ACCTCGGGCA TCGCCTACGT GCAGGTCCAG
                                                                         1440
 35 ATCACCGGCA AGGCCTCGCA TGCCGGCGCC GCGCCCGAGC TGGGCCTGAA CGCGCTGGTC
                                                                         1500
    GAGGCTTCCG ACCTCGTGCT GCGCACGATG AACATCGACG ACAAGGCGAA GAACCTGCGC
                                                                         1560
    TTCCAGTGGA CCATCGCCAA GGCCGGCCAG GTCTCGAACA TCATCCCCGC CAGCGCCACG
                                                                         1620
    CTGAACGCCG ACGTGCGCTA CGCGCGCAAC GAGGACTTCG ACGCCGCCAT GAAGACGCTG
                                                                         1680
    GAAGAGCGCG CGCAGCAGAA GAAGCTGCCC GAGGCCGACG TGAAGGTGAT CGTCACGCGC
                                                                         1740
 40 ggccgcccgg ccttcaatgc cggcgaaggc ggcaagaagc tggtcgacaa ggcggtggcc
                                                                         1800
    TACTACAAGG AAGCCGGCGG CACGCTGGGC GTGGAAGAGC GCACCGGCGG CGGCACCGAC
                                                                         186C
    GCGGCCTACG CCGCGCTCTC AGGCAAGCCA GTGATCGAGA GCCTGGGCCT GCCGGGCTTC
                                                                         1920
    GGCTACCACA GCGACAAGGC CGAGTACGTG GACATCAGCG CGATTCCGCG CCGCCTGTAC
                                                                         1980
    ATGGCTGCGC GCCTGATCAT GGATCTGGGC GCCGGCAAG
                                                                         2019
 45
```

(2) INFORMATION FOR SEQ ID NO: 32:

(i) SEQUENCE CHARACTERISTICS:

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						3 am		acio	is							
						o ac										
		(C	) ST	RAND	EDNE	SS:	sing	le								
_		(D	) TO	POLO	GY:	line	ar									
5	(ii)	MOL	ECUL	E TY	PE:	prot	ein									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 32	:					
	Met	Lys	Leu	Trp	Leu	Asn	Trp	lle	Phe	Leu	. Val	Thr	Leu	Leu	Asn	Gly
	1				5					10					15	•
10	Ile	Gln	Cys	Gln	Val	Gln	Leu	Gln	Gln	Pro	Gly	Ala	Glu	Leu	Val	Lys
				20					25					30		-
	Pro	Gly	Ala	Ser	Val	Gln	Leu	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Thr	Phe
			35					40					45	•		
	Thr	Gly	Tyr	Trp	Ile	His	Trp	Val	Lys	Gln	Arg	Pro	Glv	Gln	Glv	Leu
15		50					55					60	_		•	
	Glu	Trp	Ile	Gl.y	Glu	Val	Asn	Pro	Ser	Thr	Gly	Arg	Ser	asa	Tvr	Asn
	65					70					75	-		•		80
	Glu	Lys	Phe	Lys	Asn	Lys	Ala	Thr	Leu	Thr	Val	Asp	Lvs	Ser	Ser	
					85					90		-	-		95	
20	Thr	Ala	Tyr	Met	Gln	Leu	Ser	Ser	Leu	Thr	Ser	Glu	Asp	Ser		Val
				100					105				•	110		
	Tyr	Tyr	Cys	Ala	Arg	Glu	Arg	Ala	Tyr	Gly	Tvr	Asp	Asp		Met	Asp
	•		115					120		-	•	•	125			
	Tyr	Trp	Gly	Gln	Gly	Thr	Thr	Val	Thr	Val	Ser	Ser		Glv	Glv	Glv
25		130					135					140		,	,	1
	Ser	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Glv	Glv	Ser		Ile	Glu	Leu	Ser
	145					150	-	_	•	•	155				200	160
	Gln	Ser	Pro	Ser	Ser	Leu	Ala	Val	Ser	Ala		Glu	Lvs	Val	Thr	
					165					170	•		-2-		175	
30	Ser	Cys	Lys	Ser	Ser	Gln	Ser	Leu	Leu	Asn	Ser	Ara	Thr	Ara		Asn
				180					185			,		190	-,-	
	Tyr	Leu	Ala	Trp	Tyr	Gln	Gln	Arg	Pro	Gly	Gln	Ser	Pro		Leu	Leu
			195					200		•		•	205	-,-		
	Ile	Tyr	Trp	Ala	Ser	Thr	Arg	Thr	Ser	Glv	Val	Pro		Ara	Phe	Th r
35		210					215			•		220				
	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr		Ser	Ser	Val	Gln
	225				•	230	•				235					240
	Ala	Glu	Asp	Leu	Ala	Ile	Tvr	Tvr	Cvs	Lvs		Ser	Tyr	Thr	Len	
					245		-		-1-	250				••••	255	y
40	Thr	Phe	Gly	Gly	Gly	Thr	Lvs	Leu	Glu		Lvs	Ara	Glv	Glv		G1 v
			_	260	•				265		_,	9	Cly	270	Cly	GLY
	Ser	Gly	Glv		Glv	Ser	Glv	Glv		Glv	Ser	Gln	Tve		Λen	N c n
		•	275	,	1			280	Cly	O1y	261	GIM	285	Arg	wsb	ASII
	Val	Leu		Gln	Ala	Ala	Thr		Glu	Gla	Pro	Δ1-		T) ~	T	ጥሎ ~
45		290					295	برد	JIU	9111	rto		val	116	тÀЗ	ınr
	Leu		Lvs	Len	Val	Asn		Glu	Thr	C1 v	Th ~	300	7	21-	C1	Clar
	305		-,-			310		JIU	+ + 1 L	GIÀ		отА	ASP	WIG	GIU	
											315					320

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	Ile	Ala	Ala	Ala	Gly 325		Phe	Leu	Glu	Ala 330		Leu	Lys	Asn		
	Pho	Thr	V = 1	Th.				0 -							335	
	rne	1111	Agi	340		ser	Lys	Ser	345		Leu	Val	Val	Gly 350	Asp	Asn
5	Ile	Val	Gly 355		Ile	Lys	Gly	Arg 360		Gly	Lys	Asn	Leu 365	Leu	Leu	Met
	Ser	His 370	Met	Asp	Thr	Val	Tyr 375		Lys	Gly	Ile	Leu 380	Ala	Lys	Ala	Pro
	Phe	Arg	Val	Glu	Glv	Asp	Lvs	Ala	Tur	GLV	Pro			Ala	700	7.00
10	385				-	390			- 3 -	,	395				nsp	400
	Lys	Gly	Gly	Asn		Val	Ile	Leu	His	Thr			Leu	Leu	Lys	
					405					410					415	
	Tyr	Gly	Val	Arg 420	Asp	Tyr	Gly	Thr	Ile 425	Thr	Val	Leu	Phe	Asn 430	Thr	Asp
15	Glu	Glu	Lys	Gly	Ser	Phe	Gly	Ser	Arg	Asp	Leu	Ile	Gln	Glu	Glu	Δla
			435				-	440				•	445		010	nia
	Lys	Leu	Ala	Asp	Tyr	Val	Leu		Phe	Glu	Pro	Thr		Ala	Gly	ħ e n
		450		•	-		455					460	001	714	Gry	vah
	Glu	Lys	Leu	Ser	Leu	Glv		Ser	Glv	Πle	Δla		Val	Ġln	77 a 1	C1-
20	465	_				470			1		475		•	OIII	441	480
	Ile	Thr	Gly	Lvs	Ala		His	Ala	Glv	Δla			Glu	Leu	C)	
			-	•	485					490	7124		GIU	neu	495	var
	Asn	Ala	Leu	Val		Ala	Ser	Δsn	T.au		Len	0.50	Th.	Met		<b>T</b> 1-
				500		-,	501	, mp	505	447	Den	ALG	1111	510	ASII	TTE
25	Asp	asp	Lvs		Lvs	Asn	Len	Ara		Gla	Trn	Th-	710	Ala	T	n 1 -
	- 4		515		-, -		200	520	rne	GIII	пр	1111	525	ALG	гуя	AIS
	Glv	Gln		Ser	Δen	Tla	Tlo		212	S = =	214	m		Asn		_
		530					535	110	nra	ser	AIG		rea	ASI	Ala	Asp
	Val		Tvr	Δla	Δra	Aen		7.50	Dha	200	71-	540	M	Lys	<b></b>	
30	545		-,-		9	550	Giu	rap	riie	Map	555	WIG	met	Lys	The	
		Glo	Ara	Ala	Gln		Luc	1	T 0	Dwa		21-	•	Val	_	560
			,		565	OII.	шуз	Lys	neu	570	GIU	AId	ASP	vai		vaı
	Ile	Val	Thr	Ara		Ara	Pro	Δla	Pho		71-	C1	C1	Gly	575	•
				580	or y	****	110	VIG	585	HSII	MIG	GIA	GIU		GIÀ	гЛз
35	Lvs	Leu	Val		ĭ.vs	Δla	Va1	פומ		T	T	C1	n1.	590 Gly	G1	m).
	-, -		595	лэр	Dys	ura	VQI	600	ıyı	Tyr	гуз	GIU	605	GIÀ	Gly	Thr
	Leu	Gly	Val	Glu	Glu	Arg	Thr		Glv	Glv	Thr	Asp		Ala	ጥህጉ	Δla
		610				,	615	1	1	,		620			- , -	nia
	Ala	Leu	Ser	Gly	Lys	Pro	Val	Ile	Glu	Ser	Leu	Gly	Leu	Pro	Gly	Phe
40	625					630					635	-			•	640
	Gly	Tyr	His	Ser	Asp	Lys	Ala	Glu	Tyr	Val	Asp	Ile	Ser	Ala	Ile	
					645					650	•				655	
	Arg	Arg	Leu	Tyr	Met	Ala	Ala	Arg	Leu	Ile	Met	Asp	Leu	Gly		Glv
				660				-	665					670		,
45	Lys													-		

	(2) INFORMATION FOR SEQ ID NO: 33:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 37 base pairs	
	(B) TYPE: nucleic acid	
5		
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:	
10	GGGCGCCGGC AAGTGATAAA ATTCCTCGAG GAGCTCC	37
		3,
	(2) INFORMATION FOR SEQ ID NO: 34:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 19 base pairs	
15	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:	
20		
	CGCCACCTCT GACTTGAGC	19
	·	
	(2) INFORMATION FOR SEQ ID NO: 35:	
25	(i) SEQUENCE CHARACTERISTICS:	
25	(A) LENGTH: 37 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20	(ii) MOLECULE TYPE: other nucleic acid	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:	
	CCN COMPANY OF THE PARTY OF THE	
	GGAGCTCCTC GAGGAATTTT ATCACTTGCC GGCGCCC	37
	(2) INFORMATION FOR SEQ ID NO: 36:	
35	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 19 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 36:	
	godined backtriton, SEQ ID NO: 36:	
	GCTGAACGCC GACGTGCGC	
		19
15	(2) INFORMATION FOR SEQ ID NO: 37:	
	(i) SEQUENCE CHARACTERISTICS.	

(A) LENGTH: 2025 base pairs

```
(B) TYPE: nucleic acid(C) STRANDEDNESS: single
```

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

						TCAGTGTCAG	60
						GCAGCTGTCC	120
	TGCAAGGCTT	CTGGCTACAC	CTTCACCGGC	TACTGGATAC	ACTGGGTGAA	GCAGAGGCCT	180
10	GGACAAGGCC	TTGAGTGGAT	TGGAGAGGTT	AATCCTAGTA	CCGGTCGTTC	TGACTACAAT	240
						AGCCTACATG	300
						AGAGAGGCC	360
						CGTCTCCTCA	420
	GGTGGCGGTG	GCTCGGGCGG	TGGTGGGTCG	GGTGGCGGCG	GATCTGACAT	TGAGCTCTCA	480
15	CAGTCTCCAT	CCTCCCTGGC	TGTGTCAGCA	GGAGAGAAGG	TCACCATGAG	CTGCAAATCC	540
						CCAGCAGAGA	600
						TGGGGTCCCT	660
						CAGTGTGCAG	720
•	GCTGAAGACC	TGGCAATTTA	TTACTGCAAG	CAATCTTATA	CTCTTCGGAC	GTTCGGTGGA	780
20	GGCACCAAGC	TCGAGATCAA	ACGGGGCGGT	GGTGGCTCCG	GAGGTGGCGG	TAGCGGTGGC	840
		AGAAGCGCGA					900
		CGCTGGAGAA					960
						CACGGTCACG	1020
		CGGCCGGCCT					1080
25	GGCGGCAAGA	ACCTGCTGCT	GATGTCGCAC	ATGGACACCG	TCTACCTCAA	GGGCATTCTC	1140
		CGTTCCGCGT					1200
		ACGCGGTCAT					1260
		CCATCACCGT					1320
• •	CGCGACCTGA	TCCAGGAAGA	AGCCAAGCTG	GCCGACTACG	TGCTCTCCTT	CGAGCCCACC	1380
30	AGCGCAGGCG	ACGAAAAACT	CTCGCTGGGC	ACCTCGGGCA	TCGCCTACGT	GCAGGTCCAG	1440
		AGGCCTCGCA					1500
		ACCTCGTGCT					1560
		CCATCGCCAA					1620
2.5	CTGAACGCCG	ACGTGCGCTA	CGCGCGCAAC	GAGGACTTCG	ACGCCGCCAT	GAAGACGCTG	1680
33	GAAGAGCGCG	CGCAGCAGAA	GAAGCTGCCC	GAGGCCGACG	TGAAGGTGAT	CGTCACGCGC	1740
		CCTTCAATGC					1800
		AAGCCGGCGG					1860
		CCGCGCTCTC					1920
40	GGCTACCACA	GCGACAAGGC	CGAGTACGTG	GACATCAGCG	CGATTCCGCG	CCGCCTGTAC	1980
40	ATGGCTGCGC	GCCTGATCAT	GGATCTGGGC	GCCGGCAAGT	GATAA		2025

## (2) INFORMATION FOR SEQ ID NO: 38:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 288 amino acids

45 (B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

	(	ii)	MOL	ECUL	E TY	PE:	prot	ein									
	(:	xi)	SEQ	UENC	E DE	SCRI	PTIC	N: S	EQ I	ои д	: 38	:					
5			Lys	Tyr	Leu	Leu	Pro	Thr	Ala	Ala	Ala	Gly	Leu	Leu	Leu	Leu	Ala
5		1	C1 .			5					10					15	
	•	нта	GID	Pro		Met	Ala	Gln	Val		Leu	Gln	Gln	Pro	Gly	Ala	Glı
	,	l an	1723	T 110	20	<b>61</b>		_		25	_				30		
	•	beu	Val	35	PIO	Gly	Ата	Ser		Gln	Leu	Ser	Cys		Ala	Ser	Gl
10		rvr	Thr		<b>ም</b> ኮ ድ	Gly	Ψ	T	40	***	<b></b>		_	45			
		- , -	50		1111	Gly	ıyı	55	116	піѕ	Trp	vaı	Lys 60	Gin	Arg	Pro	Gly
	(	Sln	Gly	Leu	Glu	Trp	Ile		Glu	Val	Agn	Pro		Thr	G)	λ	c
		65	-				70	<b>U</b> _J.		***	non	75	361	1111	GIY	Arg	
	2	Asp	Tyr	Asn	Glu	Lys	Phe	Lys	Asn	Lvs	Ala		Leu	Thr	Val	Asn	80
15						85		-			90			••••	•	95	Lys
	5	Ser	Ser	Thr	Thr	Ala	Tyr	Met	Gln	Leu	Ser	Ser	Leu	Thr	Ser		Asr
					100					105					110		٠
	5	Ser	Ala	Val	Tyr	Tyr	Cys	Ala	Arg	Glu	Arg	Ala	Tyr	Gly	Tyr	Asp	Asp
20				115					120					125			
20	F	Ala		Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Thr	Val	Thr	Val	Ser	Ser	Gly
	_		130	-				135					140				
			Gly	Gly	Ser	Gly		Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Asp	Ile
		.45	•	•	٠.	_	150		-			155					160
25	٠	, Lu	Leu	ser	GID	Ser	Pro	Ser	Ser	Leu		Val	Ser	Ala	Gly	Glu	Lys
	T.	la l	Thr	Mor	S = =	165	T	0	•	۵.	170	_				175	
	•	aı	1111	Mec	180	Суз	гÀз	ser	Ser		Ser	Leu	Leu	Asn		Arg	Thr
	A	ıra	Lvs	Asn		Leu	Δla	Trn	Tur	185	C1 =	<b>7</b>	0	<b>01</b>	190	_	_
	•	5	-,0	195	- 7 -	Deu	ura	пр	200	GIII	GIU	Arg	Pro	205	GID	Ser	Pro
30	L	ys	Leu		Ile	Tyr	Tro	Ala		Thr	Ara	Th r	Sar		Wa 1	Dua	7
			210			•	•	215			•••9		220	GLY	401	FIO	ASP
	A	rg	Phe	Thr	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe		Leu	Thr	Tle	Ser
		25					230		-		•	235					240
	S	er	Val	Gln	Ala	Glu	Asp	Leu	Ala	Ile	Tyr	Tyr	Cys	Lys	Gln	Ser	
35						245					250					255	
	T	hr	Leu	Arg	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys	Arg	Glu
					260					265					270		
	G	ln	Lys		Ile	Ser	Glu	Glu	qeA	Leu	Asn	His	His	His	His	His	His
10				275		•			280					285			
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15						DNES			۵								
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	(i	i)				E: 0			leic	aci	d.						

- 83 -

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:
   GCCCAACCAG CCATGGCCGA GGTGCAGCTG CAGCAG
                                                                           36
 5 (2) INFORMATION FOR SEQ ID NO: 40:
         (i) SEQUENCE CHARACTERISTICS:
              (A) LENGTH: 54 base pairs
              (B) TYPE: nucleic acid
              (C) STRANDEDNESS: single
10
              (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: other nucleic acid
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:
   CGACCCACCA CCGCCCGAGC CACCGCCACC CGAGCTCACG GCGACTGAGG TTCC
15
   (2) INFORMATION FOR SEQ ID NO: 41:
        (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 54 base pairs
             (B) TYPE: nucleic acid
20
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: other nucleic acid -
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:
25 TCGGGCGGTG GTGGGTCGGG TGGCGGCGGA TCTCAGATTG TGCTCACCCA GTCT
                                                                           54
   (2) INFORMATION FOR SEQ ID NO: 42:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 24 base pairs
30
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: other nucleic acid
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:
35
   CCGTTTGATC TCGAGCTTGG TCCC
                                                                          24
   (2) INFORMATION FOR SEQ ID NO: 43:
        (i) SEQUENCE CHARACTERISTICS:
40
            (A) LENGTH: 843 base pairs
             (B) TYPE: nucleic acid
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: other nucleic acid
45
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:
```

ATGAAATACC TATTGCCTAC GGCAGCCGCT GGATTGTTAT TACTCGCTGC CCAACCAGCC

										٠.						
	ATGGCCG	IGG 1	rgcad	SCTGC	A G	CAGTO	CTGGG	GC#	AGAGO	CTTG	TGAG	GTC	AGG	GGCÇ'	rcag'	rc
	AAGTTGTC															
	CAGAGGCC															
_	GAATATGO	cc c	CGAAC	STTCC	G G	GCAZ	AGGCC	ACI	TTG	CTG	CAG	CTC	ATC	CTCC	AACA	CA
5	GCCTACCT	GC A	ACCTO	CAGCA	G C	TGAC	CATCT	GAG	GACA	CTG	CCGI	CTA	TA .	CTGT	CATG	C
	CTGATCTA															
	AGCTCGGG															
	CTCACCCA	CT C	CTCCA	GCAA	T CA	TGT	CTGCA	TCI	CCAG	GGG	AGA	GGT	CAC	CATA	ACCTO	SC
	AGTGCCAG	CT C	CAAGI	GTAA	C TI	ACAI	GCAC	TGG	TTCC	AGC	AGAA	GCC	AGG (	CACT	CTC	:c
10	AAACTCTG	GA I	TATT	'AGCA	C AT	CCAA	CCTG	GCI	TCTG	GAG	TCCC	TGC	rcg (	CTTC	GTGG	C
	AGTGGATC															
	ACTTATTA															
	GAGATCAA	AC G	GGAA	CAAA	A AC	TCAT	CTCA	GAA	GAAG	ATC	TGAA	TCAC	CA	CATO	ACCA	'C
	CAT															
15																
	(2) INFO															
	(i)	SEQ	UENC	E CH	ARAC	TERI	STIC	s:								
				ngth				acid	s							
20				PE: a												
20				RANDI			_	le								
				POLO												
				E TY												
	(X1)	SEQ	UENC	E DES	SCRI	PTIO	N: S	EQ I	D NO	: 44	:					
25	<b>M</b> -4	•	_		_	_										
2,5	мес 1	гÀ2	Tyr	Leu		Pro	Thr	Ala	Ala		Gly	Leu	Leu	Leu	Leu	Ala
		Cln	Dwa	71-	5	<b>7.</b> 2 -				10					15	
	*****	GIII	FIG	Ala 20	Met	Ald	GIU	vaı		Leu	Gln	Gln	Ser		Ala	Glu
	I.en	Val	Ara		G1.	71-	Co=	1103	25	•	0	_		30	_	
30	200	741	35	Ser	GIY	ATG	ser	40	гуз	Leu	ser	Cys		Ala	Ser	Gly
	Phe	Asn		Lys	Acn	) en	т		111.	m	12. 1		45		_	
		50		шys	nop	VOII	55	Mer	птэ	Trp	vaı	Lys	Gin	Arg	Pro	Glu
	Gln	_	Leu	Glu	Tro	Tle		Trn	710	λοο	Dro		<b>7</b>	63		<b></b>
	65					70	n.a	ııp	116	ASD	75	GIU	ASN	GIĀ	Asp	
35	Glu	Tyr	Ala	Pro	Lvs		Ara	Glv	Lve	Δla	. –	Lan	Th ∞	71-	7	80
		•			85					90		beu	LIL	MIA	95	Ser
	Ser	Ser	Asn	Thr	Ala	Tvr	Leu				Ser	ī.en	Thr	Sor		λcn
				100		-			105					110	Giu	nsp
	· Thr	Ala	Val	Tyr	Tyr	Cys	His	Val		Ile	Tvr	Ala	Glv		Len	Δla
40			115		_	-		120					125	-,-		
	Met	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Ser	Val	Ala	Val		Ser	GIV	ดาง
		130					135					140			,	,
	Gly	Gly	Ser	Gly-	Gly	Gly	Gly	Ser	Gly	Glv	Glv		Ser	Gln	Ile	Val
	145					150	-		-	•	155	-3				160
45	Leu	Thr	Gln	Ser	Pro	Ala	Ile	Met	Ser	Ala		Pro	Glv	Glu	Lys	
					165					170			•		175	_
	Thr	Ile	Thr	Cys	Ser	Ala	Ser	Ser	Ser	Val	Thr	Tyr	Met	His		Phe

				180					185					190			
	Gln	Gln	Lys	Pro	Gly	Thr	Ser	Pro	Lys	Leu	Trp	Ile	Tyr	Ser	Thr	Ser	
			195					200					205			•	
	Asn	Leu	Ala	Ser	Gly	Val	Pro	Ala	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	
5		210					215					220				•	
	Thr	Ser	Tyr	Ser	Leu	Thr	Ile	Ser	Arg	Met	Glu	Ala	Glu	Asp	Ala	Ala	
	225					230					235			-		240	
	Thr	Tyr	Tyr	Cys	Gln	Gln	Arg	Ser	Thr	Tyr	Pro	Leu	Thr	Phe	Glv	Ala	
					245					250					255		
10	Gly	Thr	Lys	Leu	Glu	Ile	Lys	Arg	Glu	Gln	Lys	Leu	Ile	Ser	Glu	Glu	
				260					265		_			270			
	Asp	Leu	Asn	His	His	His	His	His	His								
			275					280									
15	(2) INFO	RMATI	ON I	FOR S	SEQ 1	D NO	): 45	5:									
	(i)	SEQU	DENCE	E CHA	ARACI	ERIS	STICS	3:									
		(A)	LEN	GTH:	72	base	pai	rs									
		(B)	TY	PE: 1	nucle	eic a	cid										
		(C)	STE	RANDE	EDNES	SS: s	ing	e									
20		(D)	TO	OLOC	Y: 1	.inea	ır										
	(ii)	MOLE	CULE	TYP	E: c	ther	nuc	:leid	aci	.d							
	(xi)	SEQU	JENCE	DES	CRIF	4OI T	: SE	Q II	NO:	45:							
	TCGAGATC			AACA	AAA A	CTCF	TCT	CAG	AGAZ	GA 1	CTG	ATC	C CA	CCAT	CACC	:	60
25	ACCATTAA	TG AG	•														72
	(2) INFO	RMATI	ON E	FOR S	EQ I	D NC	: 46	:									
						סדסיםי								•			
	(i)	SEQU	JENCE	CHA	IRACT	ERIS	TICS	:						•			
20	(i)	SEQU (A)	LEN	GTH:	72	base	pai							•			
30	(i)	SEQU (A) (B)	LEN TYE	NGTH: PE: n	72 ucle	base	pai	rs									
30	(i)	SEQU (A) (B)	LEN TYE	GTH:	72 ucle	base	pai	rs									
30		(A) (B) (C) (D)	LEN TYE STF	NGTH: PE: n RANDE POLOG	72 ucle DNES Y: 1	base ic a S: s inea	pai cid ingl	rs									
30	(ii)	SEQU (A) (B) (C) (D) MOLE	LEN TYE STF TOE	OGTH: PE: n RANDE POLOG TYP	72 Tucle DNES Y: 1 PE: 0	base ic a S: s inea	pai cid ingl	rs e :leic									
	(ii)	(A) (B) (C) (D)	LEN TYE STF TOE	OGTH: PE: n RANDE POLOG TYP	72 Tucle DNES Y: 1 PE: 0	base ic a S: s inea	pai cid ingl	rs e :leic									
30	(ii) (xi)	SEQU (A) (B) (C) (D) MOLE SEQU	LENCE	NGTH: PE: n RANDE POLOG C TYPE C DES	72 SUCLE SUNCE SY: 1 PE: 0	base ic a S: s inea ther	e pai cid singl r nuc	e e eleic	NO:	46:							
	(ii) (xi)	SEQU (A) (B) (C) (D) MOLE SEQU	LEN TYE STF TOE CULE JENCE	NGTH: PE: n RANDE POLOG C TYPE C DES	72 SUCLE SUNES SY: 1 PE: 0	base ic a S: s inea ther	e pai cid singl r nuc	e e eleic	NO:	46:	ттст	'GAGA	T GA	G <b>T</b> TT	· TTGT		60
	(ii) (xi)	SEQU (A) (B) (C) (D) MOLE SEQU	LEN TYE STF TOE CULE JENCE	NGTH: PE: n RANDE POLOG C TYPE C DES	72 SUCLE SUNES SY: 1 PE: 0	base ic a S: s inea ther	e pai cid singl r nuc	e e eleic	NO:	46:	ттст	'GAGA	T GA	G <b>T</b> TT	· TTGT		60 72
	(ii) (xi) AATTCTCA' TCCCGTTT	SEQU (A) (B) (C) (D) MOLE SEQU TT AA	LEN TYE STF TOE CCULE VENCE	NGTH: PE: n RANDE POLOG C TYF C DES	72 Aucle EDNES SY: 1 PE: 0 SCRIP	base ic a S: s inea ther TION	e pai cid ingl r nuc : SE	e eleic Q II	NO:	46:	ттст	'GAGA	T GA	GTTT	· TTGT		
35	(ii) (xi)  AATTCTCA TCCCGTTTC	SEQUAL (A) (B) (C) (D) MOLE SEQUAL (C) (TT AA GA TO	LEN TYE STF TOE CULE JENCE	NGTH: PE: n RANDE POLOG TYE DES PGGTG	72 Aucle EDNES SY: 1 PE: 0 ECRIP	base ic a S: s inea ther TION	e pai cid ingl r nuc e nuc e: SE	e eleic Q II TTCA	NO:	46:	ттст	GAGA	T GA	GTTT	· TTGT		
	(ii) (xi)  AATTCTCA TCCCGTTTC	SEQUENT AAGA TO SEQUENTE AAGA TO SEQUENT	LENCE	NGTH: PE: n RANDE POLOG C TYP C DES CGGTG	72 Aucle DNES Y: 1 PE: 0 CRIP ATG	base ic a S: s inea ther TION GTGG	e pai cid ingl nuc : SE TGA	e eleic Q II TTCA	NO:	46:	TTCT	GAGA	T GA	GTTT	TTGT		
35	(ii) (xi)  AATTCTCA TCCCGTTTC	SEQUENT AAGA TO SEQUENT AAGA T	LEN TYE STF TOE CULE JENCE ATGGI CON F	NGTH: PE: R RANDE POLOG C TYE C DES CGGTG TOR S C CHA	72 SUCLE SUC	base ic a S: s inea ther TION GTGG	e pai cid singl r nuce: SE TGA	e eleic Q II TTCA	NO:	46:	TTCT	GAGA	T GA	GTTT	TTGT		
35	(ii) (xi)  AATTCTCA TCCCGTTTC	SEQUUAN (A) (B) (C) (D) MOLE SEQUUAN (A) (B) (B)	LEN TYE STF TOE CCULE HENCE TGGT : CON F LENCE LEN TYF	GTH: RANDE POLOG TYP DES DES CGGTG COR S CHA	72  Taucle CDNES  TY: 1  EE: 0  CCRIF  ATG  RACT  864  ucle	base ic a S: s inea ther TION GTGG D NO EERIS bas ic a	e pai cid ingl r nuc e: SE TTGA TTICS e pa cid	rs e e Cleic Q IC TTCA	NO:	46:	TTCI	GAGA	T GA	GTTT	· TTGT		
35	(ii) (xi)  AATTCTCA TCCCGTTTC	SEQUE (A) (B) (C) (D) MOLE SEQUE TTT AA TC SEQUE (A) (B) (C)	TYE STF TOE CCULE VENCE TYGGT CON F ENCE LEN TYF STR	NGTH: PE: r RANDE POLOG TYP DES CGGTG COR S CHA HIGTH: R RANDE	72  Taucle CDNES  The control	base ic a S: s inea ther TION GTGG D NO ERIS bas ic a S: s	e pai icid ingl ingl r nuc e se e pa cid ingl	rs e e Cleic Q IC TTCA	NO:	46:	ттст	GAGA	T GA	GTTT	TTGT		
35	(ii) (xi) AATTCTCA' TCCCGTTT' (2) INFO	SEQUE (A) (B) (C) (D) MOLE SEQUE (B) (B) (C) (D) (D)	LEN TYE STF TOE CCULE ENCE TYE CON F ENCE LEN TYE STR TOE	GGTH: RANDE POLOG TYP TOES TGGTG TGG	72 Aucle CDNES Y: 1 CCRIF CCRIF  EQ I RACT 864 ucle DNES Y: 1	base ic a S: s inea ther GTGG  D NO ERIS bas ic a S: s inea	e pai ccid ingl r : nuc e: SE TTGA TTICS e pa ccid ingl	e e cleico con control con control con	NO:	46:	TTCT	GAGA	T GA	GTTT	· TTGT ·		
35	(ii) (xi)  AATTCTCA TCCCGTTT( (2) INFO( (i))	SEQUE (A) (B) (C) (D) MOLE SEQUE TTT AA TC SEQUE (A) (B) (C)	LEN TYE TOE CCULE ENCE  CON F ENCE LEN TYF STR TOF CCULE	HIGTH: PE: r RANDE POLOG TYP TO TYP TO THE HIGHER H	72  Aucle CDNES  Y: 1  EE: 0  CCRIF  EQ I  RACT  864  ucle DNES  Y: 1  E: 0	base ic a ic a ic a ic a ic a ic a inea ic her  TION  GTGG  D NO  ERIS bas ic a S: s inea ther	e pai ccid ingl r nuc e se ccid ingl r nuc	e e e e e e e e e e e e e e e e e e e	NO:	46: TT C	TTCT	GAGA	T GA	GTTT	· TTGT ·		

	ATGAAATACC TATTGCCTAC GGCAGCCGCT GGATTGTTAT TACTCGCTGC CCAACCAGCC	60
	ATGGCCCAGG TCCAACTGCA GCAGCCTGGG GCTGAACTGG TGAAGCCTGG GGCTTCAGTG	120
	CAGCTGTCCT GCAAGGCTTC TGGCTACACC TTCACCGGCT ACTGGATACA CTGGGTGAAG	180
_	CAGAGGCCTG GACAAGGCCT TGAGTGGATT GGAGAGGTTA ATCCTAGTAC CGGTCGTTCT	240
5	GACTACAATG AGAAGTTCAA GAACAAGGCC ACACTGACTG TAGACAAATC CTCCACCACA	300
	GCCTACATGC AACTCAGCAG CCTGACATCT GAGGACTCTG CGGTCTATTA CTGTGCAAGA	360
	GAGAGGGCCT ATGGTTACGA CGATGCTATG GACTACTGGG GCCAAGGGAC CACGGTCACC	420
	GTCTCCTCAG GTGGCGGTGG CTCGGGCGGT GGTGGGTCGG GTGGCGGCGG ATCTGACATT	480
	GAGCTCTCAC AGTCTCCATC CTCCCTGGCT GTGTCAGCAG GAGAGAAGGT CACCATGAGC	540
10	TGCAAATCCA GTCAGAGTCT CCTCAACAGT AGAACCCGAA AGAACTACTT GGCTTGGTAC	600
	CAGCAGAGAC CAGGGCAGTC TCCTAAACTG CTGATCTATT GGGCATCCAC TAGGACATCT	660
	GGGGTCCCTG ATCGCTTCAC AGGCAGTGGA TCTGGGACAG ATTTCACTCT CACCATCAGC	720
	AGTGTGCAGG CTGAAGACCT GGCAATTTAT TACTGCAAGC AATCTTATAC TCTTCGGACG	780
	TTCGGTGGAG GCACCAAGCT CGAGATCAAA CGGGAACAAA AACTCATCTC AGAAGAAGAT	840
15	CTGAATCACC ACCATCACCA CCAT	864
	(2) INFORMATION FOR SEQ ID NO: 48:	
	(i) SEQUENCE CHARACTERISTICS:	
20	(A) LENGTH: 34 base pairs	
20	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:	
23	A A COMMOCA A DEPOS CHOMAS A COMMON A C	
	AAGCTTGGAA TTCAGTGTGA GGTGCAGCTG CAGC	34
	(2) INFORMATION FOR SEQ ID NO: 49:	
	(i) SEQUENCE CHARACTERISTICS:	
30	(A) LENGTH: 45 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:	
	The second of the second secon	
	CGCCACCTCC GGAGCCACCA CCGCCCCGTT TGATCTCGAG CTTGG	4.5
	Tonicion Cird	45
	(2) INFORMATION FOR SEQ ID NO: 50:	
40	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 1998 base pairs	•
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
45	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50:	

```
ATGAAGTTGT GGCTGAACTG GATTTTCCTT GTAACACTTT TAAATGGAAT TCAGTGTGAG
                                                                           60
   GTGCAGCTGC AGCAGTCTGG GGCAGAGCTT GTGAGGTCAG GGGCCTCAGT CAAGTTGTCC
                                                                          120
   TGCACAGCTT CTGGCTTCAA CATTAAAGAC AACTATATGC ACTGGGTGAA GCAGAGGCCT
                                                                          180
   GAACAGGGCC TGGAGTGGAT TGCATGGATT GATCCTGAGA ATGGTGATAC TGAATATGCC
                                                                          240
 5 CCGAAGTTCC GGGGCAAGGC CACTTTGACT GCAGACTCAT CCTCCAACAC AGCCTACCTG
                                                                          300
   CACCTCAGCA GCCTGACATC TGAGGACACT GCCGTCTATT ACTGTCATGT CCTGATCTAT
                                                                          360
   GCTGGTTATT TGGCTATGGA CTACTGGGGT CAAGGAACCT CAGTCGCCGT GAGCTCGGGT
                                                                          420
   GGCGGTGGCT CGGGCGGTGG TGGGTCGGGT GGCGGCGGAT CTCAGATTGT GCTCACCCAG
                                                                          480
   TCTCCAGCAA TCATGTCTGC ATCTCCAGGG GAGAAGGTCA CCATAACCTG CAGTGCCAGC
                                                                          540
10 TCAAGTGTAA CTTACATGCA CTGGTTCCAG CAGAAGCCAG GCACTTCTCC CAAACTCTGG
                                                                          600
   ATTTATAGCA CATCCAACCT GGCTTCTGGA GTCCCTGCTC GCTTCAGTGG CAGTGGATCT
                                                                          660
   GGGACCTCTT ACTCTCTCAC AATCAGCCGA ATGGAGGCTG AAGATGCTGC CACTTATTAC
                                                                          720
   TGCCAGCAAA GGAGTACTTA CCCGCTCACG TTCGGTGCTG GGACCAAGCT CGAGATCAAA
                                                                         780
   CGGGGCGGTG GTGGCTCCGG AGGTGGCGGT AGCGGTGGCG GGGGTTCCCA GAAGCGCGAC
                                                                         840
15 AACGTGCTGT TCCAGGCAGC TACCGACGAG CAGCCGGCCG TGATCAAGAC GCTGGAGAAG
                                                                         900
   CTGGTCAACA TCGAGACCGG CACCGGTGAC GCCGAGGGCA TCGCCGCTGC GGGCAACTTC
                                                                         960
   CTCGAGGCCG AGCTCAAGAA CCTCGGCTTC ACGGTCACGC GAAGCAAGTC GGCCGGCCTG
                                                                        1020
   GTGGTGGGCG ACAACATCGT GGGCAAGATC AAGGGCCGCG GCGGCAAGAA CCTGCTGCTG
                                                                        1080
   ATGTCGCACA TGGACACCGT CTACCTCAAG GGCATTCTCG CGAAGGCCCC GTTCCGCGTC
                                                                        1140
20 GAAGGCGACA AGGCCTACGG CCCGGGCATC GCCGACGACA AGGGCGGCAA CGCGGTCATC
                                                                        1200
   CTGCACACGC TCAAGCTGCT GAAGGAATAC GGCGTGCGCG ACTACGGCAC CATCACCGTG
                                                                        1260
   CTGTTCAACA CCGACGAGGA AAAGGGTTCC TTCGGCTCGC GCGACCTGAT CCAGGAAGAA
                                                                        1320
   GCCAAGCTGG CCGACTACGT GCTCTCCTTC GAGCCCACCA GCGCAGGCGA CGAAAAACTC
                                                                        1380
   TCGCTGGGCA CCTCGGGCAT CGCCTACGTG CAGGTCCAGA TCACCGGCAA GGCCTCGCAT
                                                                        1440
25 GCCGGCGCCG CGCCCGAGCT GGGCGTGAAC GCGCTGGTCG AGGCTTCCGA CCTCGTGCTG
                                                                        1500
   CGCACGATGA ACATCGACGA CAAGGCGAAG AACCTGCGCT TCCAGTGGAC CATCGCCAAG
   GCCGGCCAGG TCTCGAACAT CATCCCCGCC AGCGCCACGC TGAACGCCGA CGTGCGCTAC
                                                                        1620
   GCGCGCAACG AGGACTTCGA CGCCGCCATG AAGACGCTGG AAGAGCGCGC GCAGCAGAAG
                                                                        1680
   AAGCTGCCCG AGGCCGACGT GAAGGTGATC GTCACGCGCG GCCGCCCGGC CTTCAATGCC
                                                                        1740
30 ggcgaaggcg gcaagaagct ggtcgacaag gcggtggcct actacaagga agccggcggc
                                                                        1800
   ACGCTGGGCG TGGAAGAGCG CACCGGCGGC GGCACCGACG CGGCCTACGC CGCGCTCTCA
                                                                        1860
   GGCAAGCCAG TGATCGAGAG CCTGGGCCTG CCGGGCTTCG GCTACCACAG CGACAAGGCC
                                                                        1920
   GAGTACGTGG ACATCAGCGC GATTCCGCGC CGCCTGTACA TGGCTGCGCG CCTGATCATG
                                                                        1980
   GATCTGGGCG CCGGCAAG
                                                                        1998
35
   (2) INFORMATION FOR SEQ ID NO: 51:
        (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 666 amino acids
             (B) TYPE: amino acid
40
             (C) STRANDEDNESS: single
             (D) TOPOLOGY: linear
       (ii) MOLECULE TYPE: protein
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:
45
       Met Lys Leu Trp Leu Asn Trp Ile Phe Leu Val Thr Leu Leu Asn Gly
```

10

Ile Gln Cys Glu Val Gln Leu Gln Gln Ser Gly Ala Glu Leu Val Arg

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				20					25					30		
	Ser	Gly		Ser	Val	Lys	Leu		Cys	Thr	Ala	Ser	Gl	Phe	Asn	Il
			35					40					45			
5	Lys	Asp 50	Asn	Tyr	Met	His	Trp 55	Val	Lys	Gln	Arg	Pro 60	Glu	Gln	Gly	Le
	Glu 65	Trp	Ile	Ala	Trp		Asp	Pro	Glu	Asn		Asp	Thr	Glu	Tyr	
			<b>D1</b>			70 -					75					80
	Pro	ьys	Fne	Arg	61y 85	Lys	Ala	Thr	Leu	Thr 90	Ala	Asp	Ser	Ser	Ser 95	Ası
10	Thr	Ala	Tyr		His	Leu	Ser	Ser			Ser	Glu	Asp	Thr	Ala	Va.
	<b></b>	<b></b>	_	100					105					110		
	Tyr	Tyr	Cys 115	His	Val	Leu	Ile	Tyr 120		Gly	Tyr	Leu	Ala		Asp	Ту
	Trp	Gly	Gln	Gly	Thr	Ser	Val	Ala	Val	Ser	Ser	Gly	Glv	Glv	Gly	Sei
15		130					135					140		•		
	Gly	Gly	Gly	Gly	Ser	Gly	Glv	Glv	Glv	Ser	Gln			ī.eu	Thr	G1r
	145		_	_		150	-		3		155			Deu	1111	
	Ser	Pro	Ala	He	Met		A1 =	Sor	Bro	C) v			11-1	<b>(T)</b> b	Ile	160
					165	501	nia	361	FIO		GIU	гуз	Val	Inr		Thi
20	Cve	Sar	71-	80-		C	17- 1	<b>m</b> >	_	170		_			175	
	Cys	261	MIG		261	ser	vai	Inr		Met	His	Trp	Phe		Gln	Lys
	D	<b>C1</b>	<b></b>	180	_	_			185					190		
	Pro	GIY		Ser	Pro	Lys	Leu		Ile	Tyr	Ser	Thr	Ser	Asn	Leu	Ala
			195					200					205			
25	Ser	Gly	Val	Pro	Ala	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Ser	Tyr
25		210					215					220				
	Ser	Leu	Thr	Ile	Ser	Arg	Met	Glu	Ala	Glu	Asp	Ala	Ala	Thr	Tyr	Туг
	225					230					235					240
	Cys	Gln	Gln	Arg	Ser	Thr	Tyr	Pro	Leu	Thr	Phe	Gly	Ala	Gly	Thr	Lys
					245					250		_		-	255	•
30	Leu	Glu	Ile	Lys	Arg	Gly	Gly	Gly	Glv	Ser	Glv	Glv	Glv	Glv	Ser	Glu
				260	-	-		•	265		3	,	1	270	-	O.L.y
	Gly	Gly	Glv	Ser	Gln	Lvs	Arσ	Asn		Val	Leu	Dha	Gla		Ala	mh -
	-	•	275			] -	9	280			Deu	1110	285	A14	nia	1111
	Asp	Glu	_	Pro	Δla	V = 1	710		Th.	1	C1	T		17-1	Asn	٠.
35		290					295	Lys	1111	Leu	GIU		Leu	vai	ASN	TIE
	Glu		Glv	Thr	C1v	n an		C1	C1	71-		300		-1	_	
	305	****	Gry	1111	GIY		, MIG	GIU	сту	TIE		Ala	Ala	GIY	Asn	
		<b>61</b>		٠,	_	310					315					320
	Leu	GIU	Ala	Glu		Lys	Asn	Leu	Gly		Thr	Val	Thr	Arg	Ser	Lys
40					325					330					335	
40	Ser	Ala	Gly	Leu	Val	Val	Gly	Asp	Asn	Ile	Val	Gly	Lys	Ile	Lys	Gly
				340					345					350		
	Arg	Gly	Gly	Lys	nsA	Leu	Leu	Leu	Met	Ser	His	Met	Asp	Thr	Val	Tyr
			355					360					365			
	Leu	Lys	Gly	Ile	Leu	Ala	Lys	Ala	Pro	Phe	Arg	Val	Glu	Gly	Asp	Lys
45		370					375				-	380		•	•	•
	Ala	Tyr	Gly	Pro	Gly	Ile	Ala	Asp	Asp	Lys	Glv		Asn	Ala	Val	Ile
	385		_		-	390		•		-	395	-,				400

					Leu			reu	Lys	GIU	Tyr	GIA	Val	Arg	Asp	Tyr	Gly	
						405					410					415		
		Thr	Ile	Thr	Val	Leu	Phe	Asn	Thr	Asp	Glu	Glu	Lys	Gly	Ser	Phe	Gly	
					420					425					430		•	
5		Ser	Arg	Asp	Leu	Ile	Gln	Glu	Glu	Ala	Lvs	Leu	Ala	Asp	Tvr	Val	ī.en	
				435					440		,			445		141	Deu	
		Ser	Phe	Glu	Pro	Thr	Ser	Ala	Glv	Asn	Glu	Luc	T Au			Gly	m	
			450					455	01,		Olu	Dys		Jer	Leu	сту	Inr	
		Ser		Tla	Δla	Tur	T/ - 1		**- 1	G1-			460	_				
10		465		116	VTO	ıyı			vaı	GIN	ire		Gly	Lys	Ala	Ser	His	
				<b>7.1</b> -		_	470					475					480	
		ALG	GIY	MIG	мта		GIU	Leu	GIY	Val	Asn	Ala	Leu	Val	Glu	Ala	Ser	
						485					490					495		
		Asp	Leu	Val		Arg	Thr	Met	Asn	Ile	Asp	Asp	Lys	Ala	Lys	Asn	Leu	
					500			•		505					510			
15		Arg	Phe	Gln	Trp	Thr	Ile	Ala	Lys	Ala	Gly	Gln	Val	Ser	Asn	Ile	Ile	
				515					520					525				
		Pro	Ala	Ser	Ala	Thr	Leu	Asn	Ala	Asp	Val	Arq	Tyr	Ala	Ara	Asn	Glu	
			530					535					540					
		Asp	Phe	Asp	Ala	Ala	Met	Lys	Thr	Leu	Glu	Glu	Arσ	Ala	Gln	Gln	Lue	
20		545					550	•				555			0111	OIII	560	
		Lys	Leu	Pro	Glu	Ala	Asp	Val	1.vs	Val	Tla		mb ~	7	C1	Arg		
		-				565			27.5	141	570	AGI	1111	Arg	GIÀ		Pro	
		Ala	Phe	Asn	Δ1 a		C1	C1	<b>61</b>	*				_	_	575		
				11311	580	Gry	GIU	GIA	GIA		rys	Leu	Val	Asp		Ala	Val	
25		21-	m	т						585					590			
رع		ATA	TÀL		гуз	Glu	Ala	GIĀ		Thr	Leu	Gly	Val	Glu	Glu	Arg	Thr	
				595					600					605				
		GIY		Gly	Thr	Asp	Ala	Ala	Tyr	Ala	Ala	Leu	Ser	Gly	Lys	Pro	Val	
			610					615					620					
•		Ile	Glu	Ser	Leu	Gly	Leu	Pro	Gly	Phe	Gly	Tyr	His	Ser	Asp	Lys	Ala	
30		625					630					635					640	
		Glu	Tyr	Val	Asp	Ile	Ser	Ala	Ile	Pro	Arg	Arg	Leu	Tyr	Met	Ala	Ala	
				•		645					650			_		655		
		Arg	Leu	Ile	Met	Asp	Leu	Gly	Ala	Gly	Lys							
					660					665	-							
35																		
	(2)	INFOR	MATI	ON F	OR S	EO I	D NO	): 52										
			SEQU															
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		1::1				Y: 1		_										
		(ii)																
		(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	52:							
15	<b></b>		_															
IJ	GAAT	TCGCC	G CC	ACTA	TGGA	TTT	TCAA	GTG.	CAGA	TTTT	CA G	CTTC	CTGC	T AA	TCAG	TGCT		60
		TCATA								CCCA	GT C	TCCA	GCAA'	T CC	TGTC	TGCA		120
	TOTO	CAGGG	C AC	2200	mcz c	7 7 77	~ 3 ~~											

	TGGTACCAGC	AGAAGCCAGG	TTCCTCCCCC	AAATCCTGGA	TTTATGCCAC	ATCCAACCTG	240
	GCTTCTGGAG	TCCCTGCTCG	CTTCAGTGGC	AGTGGGTCTC	GGACCTCTTA	CTCTCTCACA	
						GAGTAGTAAA	360
					GGACTGTGGC		420
5	GTCTTCATCT	TCCCGCCATC	TGATGAGCAG	TTGAAATCTC	GAACTGCCTC	TGTTGTGTGC	480
					GGAAGGTGGA		540
					GCAAGGACAG		600
					AACACAAAGT		660
					GCTTCAACAG		720
10					CGACATTAAT		780
					AAACCTGGCC		840
					ATGCAAGGTC		900
					ACAACGTCTG		960
					TGCGGCCAAA		1020
15					GTTGTGAGTT		1080
					GGGCTGAAGG		1140
					ACATGCTTTA		1200
	GTCGAGGTTA	AAAAACGTCT	AGGCCCCCG	AACCACGGGG	ACGTGGTTTT	CCTTTGAAAA	1260
	ACACGATGAT	AATACCATGG	AGTTGTGGCT	GAACTGGATT	TTCCTTGTAA	CACTTTTAAA	1320
20	TGGTATCCAG	TGTGAGGTGA	AGCTGGTGGA	GTCTGGAGGA	GGCTTGGTAC	AGCCTGGGGG	1380
	TTCTCTGAGA	CTCTCCTGTG	CAACTTCTGG	GTTCACCTTC	ACTGATTACT	ACATGAACTG	1440
					TTTATTGGAA		1500
					TTCACCATCT		1560
25	ATCCCAAAGC	ATCCTCTATC	TTCAAATGAA	CACCCTGAGA	GCTGAGGACA	GTGCCACTTA	1620
23					TACTGGGGCC		1680
					TTCCCCCTGG		1740
					GTCAAGGACT		1800
					GGCGTGCACA		1860
20	TGTCCTACAG	TCCTCAGGAC	TCTACTCCCT	CAGCAGCGTC	GTGACGGTGC	CCTCCAGCAA	1920
30					CCCAGCAACA		1980
					GGTGGCGGGG		2040
					CCGGCCGTGA		2100
					GAGGGCATCG		2160
35	CGGCCTGCTC	CTCCCCCACC	TCAAGAACCT	CGGCTTCACG	GTCACGCGAA	GCAAGTCGGC	2220
55	CCTCCTCATC	TCCCACAMCG	ACATCGTGGG	CAAGATCAAG	GGCCGCGGCG	GCAAGAACCT	2280
					ATTCTCGCGA		2340
					GACGACAAGG GTGCGCGACT		2400
					GGCTCGCGCG		2460
40					CCCACCAGCG		2520
					GTCCAGATCA		2580
					CTGGTCGAGG		2640
					CTGCGCTTCC		2700
					GCCACGCTGA		2760
					ACGCTGGAAG		2820
					ACGCGCGGCC		2880
					GTGGCCTACT		2940 3000
							3000

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	CGGCGGCACG CTGGGCGTGG AAGAGCGCAC CGGCGGCGGC ACCGACGCGG CCTACGCCGC	3060
	GCTCTCAGGC AAGCCAGTGA TCGAGAGCCT GGGCCTGCCG GGCTTCGGCT ACCACAGCGA	3120
	CAAGGCCGAG TACGTGGACA TCAGCGCGAT TCCGCGCCGC CTGTACATGG CTGCGCGCCT	3180
	GATCATGGAT CTGGGCGCCG GCAAGTGATA ATCTAGA	3217
5		
	(2) INFORMATION FOR SEQ ID NO: 53:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 35 base pairs	
	(B) TYPE: nucleic acid	
10		
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 53:	
15	TGGATCTGAA GCTTAAACTA ACTCCATGGT GACCC	35
		33
	(2) INFORMATION FOR SEQ ID NO: 54:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 61 base pairs	
20	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
2.5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:	
25		
	GCCACGGATC CCGCCACCTC CGGAGCCACC ACCGCCACAA TCCCTGGGCA CAATTTTCTT	60
	G · ·	61
	/2) THEODWARTON FOR CHAIR TO MA	
30	(2) INFORMATION FOR SEQ ID NO: 55:  (i) SEQUENCE CHARACTERISTICS:	
50	(A) LENGTH: 94 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55:	
	GCCCAGGAAG CTTGGCGGTG GTGGCTCCGG AGGTGGCGGT AGCGGTGGCG GGGGTTCCCA	60
	GAAGCGCGAC AACGTGCTGT TCCAGGCAGC TACC	94
40		
	(2) INFORMATION FOR SEQ ID NO: 56:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 51 base pairs	
	(B) TYPE: nucleic acid	
45	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE, other public and	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:

	ATG	TGCC	5AAT	TCAC	GCAG	CAG	STTCT	TGCC	G CC	GCGG	CCC1	TG	TCT	rgcc	С		51
5	(2)	INI	FORM	TION	N FOR	R SEC	מד כ	NO:	57.								
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20	ATC	AGT	GCT	TCA	GTC	_	ATG	דרר		_	CDD	እ C መ	C TO TO		.0 :	CAG	
	Ile	Ser	Ala	Ser	Val	Tle	Met	Ser	non n-a	Gla	Cla	Mr.	GII	C10	. 100	CAG	99
			15					20	ALG	GIY	GIN	int		rea	Ser	GIn	
	TCT	CCA			CTG	ጥርጥ	GCA	TCT	CCN	~~~	C N C		25				
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25		30		110	Deu	361	35	Ser	PIO	CTA	GIU		Val	Thr	Met	Thr	
				ACC	<b>ጥ</b> ር አ	ACT		ĄCT	m » C	200	a.a	40					
																	195
	45		*****	Der	Ser	50	val	Thr	Tyr	TTE		Trp	Tyr	Gln	Gln		
		GCT	TCC	TCC	000		maa				55					60	
30	Pro	Gly	205	200	200	AAA	700	TGG	ATT	TAT	GCC	ACA	TCC	AAC	CTG	GCT	, 243
50	110	Gry	261	261	65	rys	ser	Trp	IIe		Ala	Thr	Ser	Asn		Ala	
	ጥርጥ	GGN	CTC	CCT		000	mm.c			70		_			75		
								AGT									291
	Der	Gry	vai		Ala	Arg	rne	Ser		Ser	Gly	Ser	Gly		Ser	Tyr	
35	TOT	CTIC	202	80					85					90			
22	207	7.00	ACA	ATC	AGC	AGA	GTG	GAG	GCT	GAA	GAT	GCT	GCC	ACT	TAT	TAC	339
	Jer.	Leu		11e	Ser	Arg	Val	Glu	Ala	Glu	Asp	Ala	Ala	Thr	Tyr	Tyr	
	mcc		95					100					105				
								CCA									387
40	Cys .		HIS	Trp	Ser	Ser		Pro	Pro	Thr	Phe	Gly	Gly	Gly	Thr	Lys	
70	cma.	110					115					120					
								GCT									435
		Glu	Ile	Lys	Arg	Ala	Asp	Ala	Ala	Pro	Thr	Val	Ser	Ile	Phe	Pro	•
	125					130					135					140	
15	CCA	TCC	AGT	GAG	CAG	TTA	ACA	TCT	GGA	GGT	GCC	TCA	GTC	GTG	TGC	TTC	483
43	Pro	Ser	Ser	Glu	Gln	Leu	Thr	Ser	Gly	Glà	Ala	Ser	Val	Val	Cys	Phe	
					145					150					155		
	TTG	AAC	AAC	TTC	<b>ጥ</b> አ ር	CCC	מממ	CAC	n m c	2 2 00							

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	Leu	Asr	a Asn	Phe	Туз	Pro	Lys	Asp	Ile	≥ Asr	ı Val	Lys	Trp	Lys	: Ile	e Asp	
				160					165					170			
																G GAC	57
5	GTA	Ser			Gln	ı Asn	Gly			ı Asr	Ser	Trp	Thr	Asp	Glr	n Asp	
,		מממ	175		3.00			180					185				
	Ser	Ture	A GAC	Ser	The	. TAC	AGC	ATG	AGC	AGC	ACC	CTC	ACG	TTC	ACC	: AAG	62
	001	190		361	1111	TAT	195		Ser	Ser	Thr			Leu	Thi	Lys	
	GAC			GAA	CGA	CAT			יי מידי	י ארר	י ייי	200		N C/T		C AAG	
10	Asp	Glu	Tyr	Glu	Arq	His	Asn	Ser	Tvr	Thr	Cve	Glu	225	The	CAC	Lys	67
	205		-		•	210			-1-	••••	215		nia	11.1	nis	220	
	ACA	TCA	ACT	TCA	ccc	ATT	GTC	AAG	AGC	TTC			AAT	GAG	TGI		72
						Ile											
					225					230					235		
15	TAA	TAAG	TAA	TC													73
	(2)	INF				SEQ											
						CHA											
20						H: 2.			aci	ds							
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25	Met	Asp	Phe	Gln	Val	Gln	Ile	Phe	Ser	Phe	Leu	Leu	Ile	Ser	Ala	Ser	
	1				5					10					15		
	Val	Ile	Met	Ser	Arg	Gly	Gln	Thr	Val	Leu	Ser	Gln	Ser	Pro	Ala	Ile	
				20					25					30			
30	Leu	Ser		Ser	Pro	Gly	Glu		Val	Thr	Met	Thr	Суѕ	Arg	Ala	Ser	
30	°	C	35	<b></b>	_			40					45				
	set	50	vai	Thr	Tyr	Ile		Trp	Tyr	Gln	Gln		Pro	Gly	Ser	Ser	
	Pro		Ser	Trn	Tlo	Tur	55	Th.	C	<b>3</b>	•	60				_	
	65	-,,,		115	116	Tyr 70	nıa	1111	ser	ASN	75	Ala	Ser	Gly	Val		
35	Ala	Arg	Phe	Ser	Gly	Ser	Glv	Ser	Glv	Thr		Tur	Sar	Lan	ጥኮ ~	80 Tlo	
		Ī			85		,		,	90	501	ıyı	361	Leu	95	TIE	
	Ser	Arg	Val	Glu	Ala	Glu	Asp	Ala	Ala		Tyr	Tyr	Cvs	Gln		Trp	
				100					105		-	_	•	110		•	
	Ser	Ser	Lys	Pro	Pro	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys	
40			115					120					125				
	Arg		Asp	Ala	Ala	Pro	Thr	Val	Ser	Ile	Phe	Pro	Pro	Ser	Ser	Glu	
		130					135					140					
		Leu	Thr	Ser	Gly	Gly	Ala	Ser	Val	Val	Cys	Phe	Leu	Asn	Asn	Phe	
45	145	D	•			150					155					160	
77	īÀĽ	LIO	ьys	Asp		Asn	Val	Lys	Trp		Ile	Asp	Gly	Ser		Arg	
	Gla	Ae-	G1 ·-	17- '	165	N	Cc.	m.	<b></b>	170		_	_		175		
	2111	11011	3 t A	٧ <b>۵</b> _	ren	Asn	ser	Trp	rnr	qzA	Gln	Asp	Ser	Lys	Asp	Ser	

- 94 -

				180	•				185					190	)		
	Thr	Tyr	Ser	Met	Ser	Ser	Thr	Leu	Thr	Leu	Thr	Lys	Asp	Glu	Tyr	Glu	
			195					200					205				
	Arg	His	Asn	Ser	Tyr	Thr	Суз	Glu	Ala	Thr	His	Lys	Thr	Ser	Thr	Ser	
5		210					215					220					
	Pro	Ile	Val	Lys	Ser	Phe	Asn	Arg	Asn	Glu	Cvs						
	225					230		_			235						
	(2)	INF	ORMA	TION	FÖR	SEO	ID.	NO.	59.								
10					CE C												
					ENGT					re							
					YPE:												
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	~mm	mm>				1				5				1	-		
25	CIT	TTA	AAT	GGT	ATC	CAG	TGT	GAG	GTG	AAG	CTG	GTG	GAG	TCT	GGA	GGA	99
23	Leu	reu		GIY	He	Gln	Cys		Val	Lys	Leu	Val	Glu	Ser	Gly	Gly ,	
			15					20					25				
					CCT												147
	GTA		Val	Gln	Pro	Gly		Ser	Leu	Arg	Leu	Ser	Cys	Ala	Thr	Ser	
20		30					35					40					
<i>5</i> 0	GGG	TTC	ACC	TTC	ACT	GAT	TAC	TAC	ATG	AAC	TGG	GTC	CCC	CAG	CCT	CCA	195
		Phe	Thr	Phe	Thr	Asp	Tyr	Tyr	Met	Asn	Trp	Val	Arg	Gln	Pro	Pro	
	45					50					55					60	
					GAG												243
25	Gly	Lys	Ala	Leu	Glu	Trp	Leu	Gly	Phe	Ile	Gly	Asn	Lys	Ala	Asn	Gly	
35					65					70			*		75		
					TAC												291
	Tyr	Thr	Thr	Glu	Tyr	Ser	Ala	Ser	Val	Lys	Gly	Arg	Phe	Thr	Ile	Ser	
				80					85					90			
40	AGA	GAT	AAA	TCC	CAA	AGC	ATC	CTC	TAT	CTT	CAA	ATG	AAC	ACC	CTG	AGA	339
40	Arg	Asp	Lys	Ser	Gln	Ser	Ile	Leu	Tyr	Leu	Gln	Met	Asn	Thr	Leu	Arg	
			95					100					105				
					GCC												387
	Ala	Glu	Asp	Ser	Ala	Thr	Tyr	Tyr	Суз	Thr	Arg	Asp	Arg	Gly	Leu	Arg	
		110					115					120					
45	TTC	TAC	TTT	GAC	TAC	TGG	GGC	CAA	GGC	ACC	ACT	CTC	ACA	GTC	TCC	TCA	435
	Phe	Tyr	Phe	Asp	Tyr	Trp	Gly	Gļn	Gly	Thr	Thr	Leu	Thr	Val	Ser	Ser	
	125					130					135					140	

- 95 -

	GCC	AAA	ACG	ACA	ccc		TI CT	CTC	m 2 4		- 05					GCT	
	Ala	Lvs	Thr	The	D = 0	Due	0	G10	TAL	- CCA	· CTC	5 GCC	CCI	GG	A TCI	GCT	483
		-,,			145		3er	val	ryz			ı Ala	Pro	G13	/ Ser	Ala	
	ccc	CAA	n Cm	220						150					155		
5	33.	CAA	MCI	AAC	. 100	ATG	GTG	ACC	CTG	GGA	TGC	CTG	GTC	AAG	GGC	TAT	531
J	Ата	GIN	Thr			Met	Val	Thr	Leu	Gly	Cys	Leu	Val	Lys	Gly	Tyr	
		•		160					165					170			
	TTC	CCT	GAG	CCA	GTG	ACA	GTG	ACC	TGG	AAC	тст	' GGA	TCT	CTG	TCC	AGC	579
	Phe	Pro	Glu	Pro	Val	Thr	Val	Thr	Trp	Asn	Ser	Gly	Ser	Leu	Ser	Ser	
			175					180					185				
10	GGT	GTG	CAC	ACC	TTC	CCA	GCT	GTC	CTG	CAG	TCT	GAC	CTC	TAC	ACT	CTG	627
	Gly	Val	His	Thr	Phe	Pro	Ala	Val	Leu	Gln	Ser	Asp	Leu	Tyr	Thr	Leu	
		190					195					200		-			
	AGC	AGC	TCA	GTG	ACT	GTC	ccc	TCC	AGC	ACC	TGG	ccc	AGC	GAG	ACC	GTC	675
						Val											0,5
15	205					210					215		-	014	• • • • • • • • • • • • • • • • • • • •	220	
	ACC	TGC	AAC	GTT	GCC	CAC	CCG	GCC	AGC	AGC			CTC	CNC	770		700
						His											723
		•			225			714	Der	230	1112	гуs	val	ASP		гуз	
	ATT	GTG	CCC	AGG		TGT	GGC	CCT	CCT		maa				235		
20	Ile	Val	Pro	Δτα	) an	Cuc	C1	GGI	GGI	000	TCC	GGA	GGT	GGC	GGT	AGC	771
				240	usp	Cys	GTĀ	GIY		GIÀ	ser	GIY	GIY		Gly	Ser	
	CCT	ccc	ccc		mcc.	C . C			245					250			
						CAG											819
	GIY	GIA		GIĀ	ser	Gln	Lys		Asp	Asn	Val	Leu	Phe	Gln	Ala	Ala	
25		~- ~	255					260					265				
23	ACC	GAC	GAG	CAG	CCG	GCC	GTG	ATC	AAG	ACG	CTG	GAG	AAG	CTG	GTC	AAC	867
	Thr		Glu	Gln	Pro	Ala	Val	Ile	Lys	Thr	Leu	Glu	Lys	Leu	Val	Asn	
		270					275					280					
						GGT											915
20	Ile	Glu	Thr	Gly	Thr	Gly	qeA	Ala	Glu	Gly	Ile	Ala	Ala	Ala	Gly	Asn	
30	285					290				•	295					300	
						CTC											. 963
	Phe	Leu	Glu	Ala	Glu	Leu	Lys	Asn	Leu	Gly	Phe	Thr	Val	Thr	Arg	Ser	
					305					310					315		
	AAG	TCG	GCC	GGC	CTG	GTG	GTG	GGC	GAC	AAC	ATC	GTG	GGC	AAG	ATC	AAG	1011
35	Lys	Ser	Ala	Gly	Leu	Val	Val	Gly	Asp	Asn	Ile	Val	Gly	Lys	Ile	Lys	
				320					325				-	330		-	
	GGC	CGC	GGC	GGC	AAG	AAC	CTG	CTG	CTG	ATG	TCG	CAC	ATG	GAC	ACC	GTC	1059
						Asn											2005
			335					340					345				
40	TAC	CTC	AAG	GGC	ATT	CTC	GCG	AAG	GCC	CCG	TTC	CGC		CAA	GGC	GAC	1107
	Tyr	Leu	Lys	Gly	Ile	Leu	Ala	Lvs	Ala	Pro	Phe	Ara	Val	Clu	C1	A.S.	1107
	_	350	-	-			355	-,-			1110	360	V (1)	GIU	GIA	АЗР	
	AAG	GCC	TAC	GGC	CCG	GGC		ccc	GNC	CAC	7 7 C		000	330			
						Gly											1155
45	365		-1-	~- y	-10	370	~~~	urq	asp	usb		GIÀ	GIA	ASN 0	ATA		
		СТС	CAC	ACC.	CTC		CT C	CTC C	3 B C	a	375					380	
						AAG											1203
	- T G	T-C II	urz	1111	ren	Lys	ren	ьeu	Lys	Glu	Tyr	Gly	Val	Ara	Asp	Tvr	

					383	,				390	)				395	i	
	GGC	ACC	: ATC	ACC	GTG	CTG	TTC	: AAC	ACC	GAC	GAC	GAZ	AAG	GG1	TCC	TTC	1251
	Gly	Thr	Ile	Thr	Val	. Leu	Phe	Asr	Thr	Asp	Glu	ı Glu	Lys	Gly	, Ser	Phe	
				400					405				_	410			
5	GGC	TCG	CĞC	GAC	CTG	ATC	CAG	GAA	GAA	GCC	AAC	сто	GCC	GAC	TAC	GTG	1299
	Gly	Ser	Arg	Asp	Leu	Ile	Gln	Glu	Glu	Ala	Lys	Leu	Ala	Asc	Tvr	Val	
			415					420			-		425		•		
	CTC	TCC	TTC	GAG	ccc	ACC	AGC	GCA	GGC	GAC	GAA	AAA	cro	тс	CTG	GGC	1347
									Gly								2017
10		430					435			_		440	•				
	ACC	TCG	GGC	ATC	GCC	TAC	GTG	CAG	GTC	AAC	ATC	ACC	GGC	AAG	GCC	TCG	1395
									Val								2000
	445					450					455		-	•		460	
	CAT	GCC	GGC	GCC	GCG	ccc	GAG	CTG	GGC	GTG	AAC	GCG	CTG	GTC	GAG		1443
15	His	Ala	Gly	Ala	Ala	Pro	Glu	Leu	Gly	Val	Asn	Ala	Leu	Val	Glu	Ala	2.15
					465					470					475		
	TCC	GAC	CTC	GTG	CTG	CGC	ACG	ATG	AAC	ATC	GAC	GAC	AAG	GCG	AAG	AAC	1491
									Asn								
				480					485				-	490	•		
20	CTG	CGC	TTC	AAC	TGG	ACC	ATC	GCC	AAG	GCC	GGC	AAC	GTC	TCG	AAC	ATC	1539
	Leu	Arg	Phe	Asn	Trp	Thr	Ile	Ala	Lys	Ala	Gly	Asn	Val	Ser	Asn	Ile	1007
			495					500			_		505				
	ATC	CCC	GCC	AGC	GCC	ACG	CTG	AAC	GCC	GAC	GTG	CGC	TAC	GCG	CGC	AAC	1587
									Ala								
25		510					515					520			_		
	GAG	GAC	TTC	GAC	GCC	GCC	ATG	AAG	ACG	CTG	GAA	GAG	CGC	GCG	CAG	CAG	1635
									Thr								
	525					530					535					540	
	AAG	AAG	CTG	CCC	GAG	GCC	GAC	GTG	AAG	GTG	ATC	GTC	ACG	CGC	GGC	CGC	1683
30	Lys	Lys	Leu	Pro	Glu	Ala	Asp	Val	Lys	Val	Ile	Val	Thr	Arg	Gly	Arg	
					545					550					555		
									GGC								1731
	Pro	Ala	Phe	Asn	Ala	Gly	Glu	Gly	Gly	Lys	Lys	Leu	Val	Asp	Lys	Ala	
~ ~				560					565					570			
35	GTG	GCC	TAC	TAC	AAG	GAA	GCC	GGC	GGC	ACG	CT.G	GGC	GTG	GAA	GAG	CGC	1779
	Val	Ala	Tyr	Tyr	Lys	Glu	Ala	Gly	Gly	Thr	Leu	Gly	Val	Glu	Glu	Arg	
			575					580					585				
									TAC								1827
40	Thr	Gly	Gly	Gly	Thr	Asp	Ala	Ala	Tyr	Ala	Ala	Leu	Ser	Gly	Lys	Pro	
40		590					595					600					
									GGC								1875
		Ile	Glu	Ser	Leu	Gly	Leu	Pro	Gly	Phe	Gly	Tyr	His	Ser	Asp	Lys	
	605					610					615					620	
15	GCC	GAG	TAC	GTG	GAC	ATC	AGC	GCG	ATT	CCG	CGC	CGC	CTG	TAC	ATG	GCT	1923
43	Ala	Glu	Tyr	Val	Asp	Ile	Ser	Ala	Ile	Pro	Arg	Arg	Leu	Tyr	Met	Ala	
					625					630					635		
	GCG	CCC	CTG	ATC	ATG	GAT	CTG	GGC	GCC	GGÇ	AAG	TGAT	'AAGA	AT T	CCTC	GAG	1974

Ala Arg Leu Ile Met Asp Leu Gly Ala Gly Lys 640 645

- (2) INFORMATION FOR SEQ ID NO: 60:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 647 amino acids
  - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- 10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:

Met Lys Leu Trp Leu Asn Trp Ile Phe Leu Val Thr Leu Leu Asn Gly 10 Ile Gln Cys Glu Val Lys Leu Val Glu Ser Gly Gly Gly Leu Val Gln 25 Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Thr Ser Gly Phe Thr Phe 40 45 Thr Asp Tyr Tyr Met Asn Trp Val Arg Gln Pro Pro Gly Lys Ala Leu 55  $20~{
m Glu}$  Trp Leu Gly Phe Ile Gly Asn Lys Ala Asn Gly Tyr Thr Thr Glu 70 Tyr Ser Ala Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Lys Ser 90 Gln Ser Ile Leu Tyr Leu Gln Met Asn Thr Leu Arg Ala Glu Asp Ser 25 100 105 Ala Thr Tyr Tyr Cys Thr Arg Asp Arg Gly Leu Arg Phe Tyr Phe Asp 115 120 Tyr Trp Gly Gln Gly Thr Thr Leu Thr Val Ser Ser Ala Lys Thr Thr 135 140 30 Pro Pro Ser Val Tyr Pro Leu Ala Pro Gly Ser Ala Ala Gln Thr Asn 150 155 Ser Met Val Thr Leu Gly Cys Leu Val Lys Gly Tyr Phe Pro Glu Pro 170 Val Thr Val Thr Trp Asn Ser Gly Ser Leu Ser Ser Gly Val His Thr 35 185 Phe Pro Ala Val Leu Glm Ser Asp Leu Tyr Thr Leu Ser Ser Ser Val 200 Thr Val Pro Ser Ser Thr Trp Pro Ser Glu Thr Val Thr Cys Asn Val 215 220 40 Ala His Pro Ala Ser Ser Thr Lys Val Asp Lys Lys Ile Val Pro Arg 230 Asp Cys Gly Gly Gly Ser Gly Gly Gly Gly Ser Gly Gly Gly 250 Ser Gln Lys Arg Asp Asn Val Leu Phe Gln Ala Ala Thr Asp Glu Gln 260 265 Pro Ala Val Ile Lys Thr Leu Glu Lys Leu Val Asn Ile Glu Thr Gly 280

	Thr	: Gl <sub>y</sub> 290		Ala	Glu	Gly	11e		Ala	a Ala	a Gl	y Asi 30		e Leu	ı Glı	ı Al
			Lys	Asr	Leu	Gly	Phe		: Va]	l Thi	r Ar			s Sei	: Ala	G1
_	305					310					31					32
3	Leu	val	. Val	. Gly	Asp 325	Asn	Ile	· Val	. Gly	Ly:		E Ly:	s Gly	y Arg	Gl <sub>y</sub> 335	
	Lys	Asn	Leu	Leu	Leu	Met	Ser	His	Met	Asr	Th	r Val	Tur	r I.a.		
				340					345					350		. 01
	Ile	Leu	Ala	Lys	Ala	Pro	Phe	Ara			. 61.	, Acr	. T.,,			
10			355					360			. 01)	, ust	365		LIYI	GI
	Pro	Glv	Ile	Ala	Asp	Asp	I.vs			, her		, tř. 1				-
		370				1.55	375		GLY	MSI	1 WIS			: ьеч	HIS	Th
	Len			Lau	Tue	Clu				•	_	380				
	385		LCu	Deu	nys	Glu	Tyr	GIY	vaı	. Arg			Gly	' Thr	Ile	Th
15			Dh -	<b>.</b>	<b>.</b>	390					395					40
13	AGI	neu	rne	ASI		Asp	GIU	GIu	Lys			Phe	Gly	Ser	Arg	As
	•				405					410					415	
	Leu	116	GIn		Glu	Ala	Lys	Leu	Ala	Asp	Tyr	Val	. Leu	Ser	Phe	G1
	_			420					425					430		
20	Pro	Thr		Ala	Gly	Asp	Glu	Lys	Leu	Ser	Leu	Gly	Thr	Ser	Gly	11
20			435					440					445			
	Ala	Tyr	Val	Gln	Val	Asn	Ile	Thr	Gly	Lys	Ala	Ser	His	Ala	Gly	Al
		450					455					460				
	Ala	Pro	Glu	Leu	Gly	Val	Asn	Ala	Leu	Val	Glu	Ala	Ser	Asp	Leu	Va.
	465					470					475			_		48
25	Leu	Arg	Thr	Met	Asn	Ile	Asp	Asp	Lys	Ala	Lvs	Asn	Leu	Ara	Phe	Ası
					485				-	490				9	495	
	Trp	Thr	Ile	Ala	Lys	Ala	Gly	Asn	Val	Ser	Asn	Tle	Tle	Pro		Sei
				500	-		-		505					510	V10	36,
	Ala	Thr	Leu	Asn	Ala	Asp	Val	Ara		Δla	Δτσ	Δen	Gly		Dho	λ
30			515			•		520	. , .	*****		N3II	525	vob	rne	ASĮ
	Ala	Ala	Met	Lvs	Thr	Leu	Ghi		7 50	A1 a	C) =	C1-		•	•	_
		530		-4-			535	014	nry	Ala	GIII		гда	гуя	ьeu	Pro
	Glu		Asp	Val	Lare	Val		37-1	mb	B	<b>C1</b>	540	_			
	545				Dy 3	550	116	AGI	Ing	Arg		Arg	Pro	Ala	Phe	
35		GIV	Glu	G1.	C1		T	•			555					560
		O <sub>L</sub> y	GIU	GIY	565	Lys	гÀ2	ren	vaı		rys	Ala	Val	Ala		Tyr
	Tue	C1	71-	<b>63</b>	-		_			570					575	
	ьys	GIU	ALG		GIÀ	Thr	Leu	Gly		Glu	Glu	Arg	Thr	Gly	Gly	Gly
	m\	_		580	_				585	,				590		
40	ınr	Asp		Ala	Tyr	Ala	Ala	Leu	Ser	Gly	Lys	Pro	Val	Ile	Glu	Ser
	_		595					600					605			
	Leu	Gly	Leu	Pro	Gly	Phe	Gly	Tyr	His	Ser	Asp	Lys	Ala	Glu	Tyr	Val
		610					615					620				
	Asp	Ile	Ser	Ala	Ile	Pro	Arg	Arg	Leu	Tyr	Met	Ala	Ala	Arg	Leu	Ile
	625					630					635					640
45	Met	Asp	Leu	Gly	Ala	Gly	Lys									
					645											

### **CLAIMS**

- A gene construct encoding a cell targeting moiety and a heterologous prodrug activating enzyme for use as a medicament in a mammalian host wherein the gene construct is capable of expressing the cell targeting moiety and heterologous prodrug activating enzyme as a conjugate within a cell in the mammalian host and a least of the conjugate within a cell in the mammalian host and a least of the cell targeting moiety and heterologous prodrug activating enzyme as
- 5 a conjugate within a cell in the mammalian host and wherein the conjugate is directed to leave the cell thereafter for selective localisation at a cell surface antigen recognised by the cell targeting moiety.
  - 2 A gene construct for use as a medicament according to claim 1 wherein the cell targeting moiety is an antibody.
- A gene construct for use as a medicament according to claim 2 wherein the antibody is an anti-CEA antibody selected from antibody A5B7 or 806.077 antibody.
  - 4 A gene construct for use as a medicament according to any preceding claim wherein the heterologous prodrug activating enzyme is a carboxypeptidase.
- 5 A gene construct for use as a medicament according to claim 4 wherein the 15 carboxypeptidase is CPG2.
  - A gene construct for use as a medicament according to claim 5 wherein the CPG2 has mutated polypeptide glycosylation sites so as to prevent or reduce glycosylation on expression in mammalian cells.
- A gene construct for use as a medicament according to any one of claims 5-6 in which the antibody-enzyme CPG2 conjugate is a fusion protein in which the enzyme is fused to the C terminus of the antibody through the heavy or light chain thereof whereby dimerisation of the encoded conjugate when expressed can take place through a dimerisation domain on CPG2.
- A gene construct for use as a medicament according to claim 7 wherein the fusion protein is formed through linking a C-terminus of an antibody Fab heavy chain to an N-terminus of a CPG2 molecule to form a Fab-CPG2 whereby two Fab-CPG2 molecules when expressed dimerise through CPG2 to form a (Fab-CPG2)<sub>2</sub> conjugate.
  - A gene construct for use as a medicament according to claim 4 wherein the carboxypeptidase is selected from [D253K]HCPB, [G251T,D253K]HCPB or
- 30 [A248S,G251T,D253K]HCPB.

- A gene construct for use as a medicament according to any preceding claim comprising a transcriptional regulatory sequence which comprises a promoter and a control element which comprises a genetic switch to control expression of the gene construct.
- A gene construct for use as a medicament according to claim 10 in which the transcriptional regulatory sequence comprises a genetic switch control element regulated by presence of tetracycline or ecdysone.
- A gene construct for use as a medicament according to claim 10 or 11 wherein thepromoter is dependent on cell type and is selected from the following promoters:
  carcinoembryonic antigen (CEA); alpha-foetoprotein (AFP); tyrosine hydroxylase; choline
  acetyl transferase; neurone specific enolase; insulin; glial fibro acidic protein; HER-2/neu; cerbB2; and N-myc.
  - A gene construct for use as a medicament according to any preceding claim which is packaged within an adenovirus for delivery to the mammalian host.
- Use of a gene construct as defined in any one of claims 1-12 for manufacture of a medicament for cancer therapy in a mammalian host.
  - A matched two component system designed for use in a mammalian host in which the components comprise:
  - (i) a first component that comprises a gene construct as defined in any one of claims 1-13 and;
- 20 (ii) a second component that comprises a prodrug which can be converted into a cytotoxic drug by the heterologous enzyme encoded by the first component.
  - A matched two component system according to claim 15 in which:
    the first component comprises a gene encoding the heterologous enzyme CPG2; and
    the second component prodrug is selected from N-(4-[N,N-bis(2-iodoethyl)amino]-
- 25 phenoxycarbonyl)-<u>L</u>-glutamic acid, <u>N</u>-(4-[<u>N,N</u>-bis(2-chloroethyl)amino]phenoxycarbonyl)-<u>L</u>-glutamic-gamma-(3,5-dicarboxy)anilide or <u>N</u>-(4-[<u>N,N</u>-bis(2chloroethyl)amino]-phenoxycarbonyl)-<u>L</u>-glutamic acid or a pharmaceutically acceptable salt thereof.
- A method for the delivery of a cytotoxic drug to a site which comprises administering to a host a first component that comprises a gene construct as defined in any one of claims 1-13; followed by administration to the host of a second component that comprises a prodrug

which can be converted into a cytotoxic drug by the heterologous enzyme encoded by the first component.

- A method according to claim 17 in which the first component comprises a gene encoding the heterologous enzyme CPG2; and the second component prodrug is selected from
- 5 <u>N</u>-(4-[<u>N,N</u>-bis(2-iodoethyl)amino]phenoxycarbonyl)-<u>L</u>-glutamic acid, <u>N</u>-(4-[<u>N,N</u>-bis(2-chloroethyl)amino]-phenoxycarbonyl)-<u>L</u>-glutamic-gamma-(3,5-dicarboxy)anilide or <u>N</u>-(4-[<u>N,N</u>-bis(2-chloroethyl)amino]-phenoxycarbonyl)-<u>L</u>-glutamic acid or a pharmaceutically acceptable salt thereof.

Figure 1



Figure 2

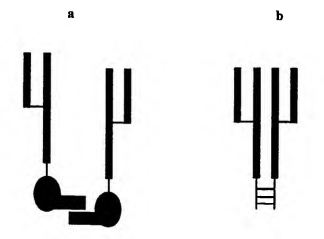


Figure 3

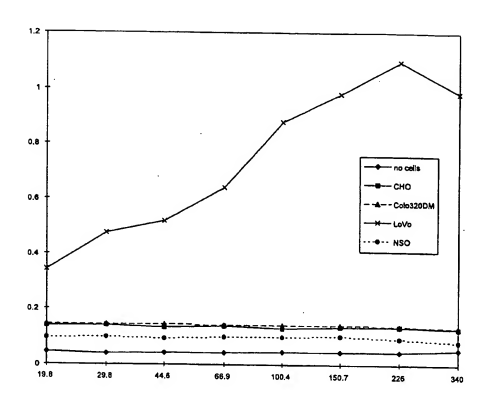


Figure 4

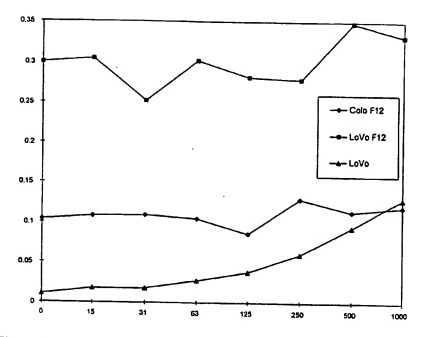
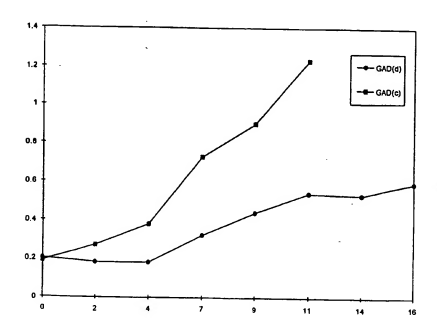


Figure 5



# **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



(CA) Turamana and District China in the	HED (	JNDER THE PATENT COOPERATION TREATY (PCT)
(51) International Patent Classification <sup>6</sup> : C12N 15/62, A61K 48/00 // C07K 16/28,	A3	(11) International Publication Number: WO 98/51787
C12N 9/48		(43) International Publication Date: 19 November 1998 (19.11.98)
(21) International Application Number: PCT/GB (22) International Filing Date: 5 May 1998 ( (30) Priority Data: 9709421.3 10 May 1997 (10.05.97)	05.05.9i G	BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPC B patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasiar patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), Europear patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR
<ul> <li>(71) Applicant (for all designated States except US): 2 LIMITED [GB/GB]; 15 Stanhope Gate, London W (GB).</li> <li>(72) Inventors; and</li> <li>(75) Inventors/Applicants (for US only): EMERY, Stephen [GB/GB]; Zeneca Pharmaceuticals, Intellectual Dept., Mereside, Alderley Park, Macclesfield, SK10 4TG (GB). BLAKEY, David, Charles [ Zeneca Pharmaceuticals, Intellectual Property Dep side, Alderley Park, Macclesfield, Cheshire Sk (GB).</li> <li>(74) Agent: GILES, Allen, Frank; Zeneca Pharmaceuticals, tual Property Dept., Mereside, Alderley Park, Mac Cheshire SK10 4TG (GB).</li> </ul>	A, Charle Proper Cheshi GB/GB ot., Men	Published  With international search report.  Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.  (88) Date of publication of the international search report:  I April 1999 (01.04.99)
MOIETY (57) Abstract  The invention provides a gene construct encoding a medicament in a mammalian host wherein the gene construct	cell targ act is ca e conju	US PRODRUG-ACTIVATING ENZYME AND A CELL TARGETING setting moiety and a heterologous prodrug activating enzyme for use as a pable of expressing the cell targeting moiety and enzyme as a conjugate gate is directed to leave the cell thereafter for selective localisation at a

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X Furth	er documents are listed in the continuation of box C.	X Patent family members are listed in	n annex.						
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mational application No.

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Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
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This international Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
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As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid specifically claims Nos.:  .
No required additional search fees were timely paid by the applicant. Consequently, this International Search Report Is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

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